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## PROVISIONAL SPECIFICATION

for the invention entitled:

"Viral variants; detection and application"

The invention is described in the following statement:

# VIRAL VARIANTS; DETECTION AND APPLICATION

### BACKGROUND OF THE INVENTION

### 5 FIELD OF THE INVENTION

The present invention relates generally to viral variants exhibiting reduced sensitivity to particular agents and/or reduced interactivity with immunological reagents. More particularly, the present invention is directed to hepatitis B virus (HBV) variants exhibiting complete or partial resistance to nucleoside or nucleotide analogs and/or reduced interactivity with antibodies to viral surface components including reduced sensitivity to these antibodies. The present invention further contemplates assays for detecting such viral variants, which assays are useful in monitoring anti-viral therapeutic regimens and in developing new or modified vaccines directed against viral agents and in particular HBV variants. The present invention also contemplates the use of the viral variants to screen for and/or develop or design agents capable of inhibiting infection, replication and/or release of the virus.

### DESCRIPTION OF THE PRIOR ART

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Bibliographic details of the publications referred to in this specification are also collected at the end of the description.

The reference to any prior art in this specification is not, and should not be taken as, an acknowledgment or any form of suggestion that that prior art forms part of the common general knowledge in any country.

Hepatitis B virus (HBV) can cause debilitating disease conditions and can lead to acute liver failure. HBV is a DNA virus which replicates *via* an RNA intermediate and utilizes reverse transcription in its replication strategy (Summers and Mason, *Cell 29*: 403-415, 1982). The HBV genome is of a complex nature having a partially double-stranded DNA

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structure with overlapping open reading frames encoding surface, core, polymerase and X genes. The complex nature of the HBV genome is represented in Figure 1. The polymerase consists of four functional regions, the terminal protein (TP), spacer, reverse transcriptase (rt) and ribonuclease (RNAse).

The polymerase gene of HBV overlaps the envelope gene, mutations in the catalytic domain of the polymerase gene can also affect the nucleotide and the deduced amino acid sequence of the envelope protein and vice versa. In particular, the genetic sequence for the neutralization domain of HBV known as the 'a' determinant, which is found within the HBsAg and located between amino acids 99 and 169, actually overlaps the major catalytic regions of the viral polymerase protein and in particular domains A and B.

The presence of an HBV DNA polymerase has led to the proposition that nucleoside or nucleotide analogs could act as effective anti-viral agents. Examples of nucleoside analogs currently being tested are penciclovir and its oral form (FCV) [Vere Hodge, Antiviral Chem Chemother 4: 67-84, 1993; Boyd et al., Antiviral Chem Chemother. 32: 358-363, 1987; Kruger et al., Hepatology 22: 219A, 1994; Main et al., J. Viral Hepatitis 3: 211-215, 1996], Lamivudine [(-)-β-2'-deoxy-3'-thiacytidine]; (3TC or LMV) [Severini et al., Antimicrobial Agents Chemother. 39: 430-435, 1995; Dienstag et al., New England J Med 333: 1657-1661, 1995]. New nucleoside or nucleotide analogs which have already 20 progressed to clinical trials include the pyrimidines Emtricitabine, ((-)-β-L-2'-3'-dideoxy-5-fluoro-3'-thiacydidine; FTC), the 5-fluoro derivative of 3TC, and Clevudine (1-(2fluoro-5-methyl-β-L-arabino-furanosyl) uracil; L-FMAU), a thymidine analog. Like 3TC, these are pyrimidine derivatives with an unnatural "L"- configuration. Several purine derivatives have also progressed to clinical trials; they include Entecavir (BMS-200, 475; 25 ETV), a carbocyclic deoxyguanosine analog, diaminopurine dioxolane (DAPD), an oral pro-drug for dioxolane guanine ((-)-β-D-2-aminopurine dioxolane; DXG) and Adefovir dipivoxil, an oral prodrug for the acyclic deoxyadenosine monophosphate nucleoside analog Adefovir (9-[phosphonyl-methoxyethyl]-adenine; PMEA). Other drugs in preclinical and clinical trials include FLG [Medivir], ACH-126,443 (L-d4C) [Archillion 30 Pharmaceuticals], ICN 2001-3 (ICN) and Racivir (RCV) [Pharmassett].

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Whilst these agents are highly effective in inhibiting HBV DNA synthesis, there is the potential for resistant mutants of HBV to emerge during long term antiviral chemotherapy. In patients on prolonged LMV therapy, key resistance mutations are selected in the rt domain within the polymerase at rtM204I/V +/- rtL180M as well as other mutations. The nomenclature used for the polymerase mutations is in accordance with that proposed by Stuyver et al., 2001, supra. LMV is a nucleoside analog that has been approved for use against chronic HBV infection. LMV is a particularly potent inhibitor of HBV replication and reduces HBV DNA titres in the sera of chronically infected patients after orthotopic liver transplantation (OLT) by inhibiting viral DNA synthesis. LMV monotherapy seems unlikely to be able to control HBV replication in the longer term. This is because emergence of LMV-resistant strains of HBV seems almost inevitable during monotherapy.

Adefovir dipivoxil (ADV: formerly, bis-pom PMEA) is an orally available prodrug of the acyclic deoxyadenosine monophosphate analog adefovir (formerly, PMEA) (Figure 2). ADV is also a potent inhibitor of HBV replication and has recently been given FDA approval for use against chronic HBV infection. Adefovir dipivoxil differs from other agents in this class in that it is a nucleotide (vs. nucleoside) analog and as such bypasses the first phosphorylation reaction during drug activation. This step is often rate-limiting. Adefovir dipivoxil has demonstrated clinical activity against both wild-type and lamivudine-resistant strains of HBV and is currently in phase III clinical Testing (Gilson et al., J Viral Hepat 6: 387-395, 1999; Perrillo et al., Hepatology 32: 129-134, 2000; Peters et al., Transplantation 68: 1912-1914, 1999; Benhamou et al., Lancet 358: 718-723, 2001). During phase II studies a 30 mg daily dose of adefovir dipivoxil resulted in a mean 4 log<sub>10</sub> decrease in viremia over 12 weeks (Heathcote et al., Hepatology 28: A620, 1998).

ADV is a substituted acyclic nucleoside phosphonate. This class of compounds also includes tenofovir disoproxil fumarate (also referred to as tenofovir DF, or tenofovir, or (TFV) or 9-R-(2-phosphonomethoxypropyl)adenine (PMPA) and is marketed as Viread by Gilead sciences).

TFV has antiviral activity against both HBV and HIV (Ying et al., J Viral Hepat. 7(2): 161-165, 2000; Ying et al., J. Viral Hepat. 7(1): 79-83, 2000; Suo et al., J Biol Chem. 273(42): 27250-27258. 1998).

5 FTC has activity against HBV and HIV (Frick et al., Antimicrob Agents Chemother 37: 2285-2292, 1993).

Nucleoside or nucleotide analog therapy may be administered as monotherapy or combination therapy where two or more nucleoside or nucleotide analogs may be administered. The nucleoside or nucleotide analogs may also be administered in combination with other antiviral agents such as interferon or hepatitis B immunoglobulin (HBIG).

There is a need to monitor for the emergence of nucleoside/nucleotide-analog- or antibodyresistant strains of HBV and to develop diagnostic protocols to detect these resistant viruses and/or to use them to screen for and/or develop or design agents having properties making them useful as anti-viral agents. Defective forms of these resistant strains or antigenic components therefrom are also proposed to be useful in the development of therapeutic vaccine compositions as are antibodies directed to viral surface components.

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### SUMMARY OF THE INVENTION

Throughout this specification, unless the context requires otherwise, the word "comprise", or variations such as "comprises" or "comprising", will be understood to imply the inclusion of a stated element or integer or group of elements or integers but not the exclusion of any other element or integer or group of elements or integers.

Nucleotide and amino acid sequences are referred to by a sequence identifier number (SEQ ID NO:). The SEQ ID NOs: correspond numerically to the sequence identifiers <400>1 (SEQ ID NO:1), <400>2 (SEQ ID NO:2), etc. A summary of the sequence identifiers is provided in Table 1. A sequence listing is provided after the claims.

Specific mutations in an amino acid sequence are represented herein as "Xaa<sub>1</sub>nXaa<sub>2</sub>" where Xaa<sub>1</sub> is the original amino acid residue before mutation, n is the residue number and Xaa<sub>2</sub> is the mutant amino acid. The abbreviation "Xaa" may be the three letter or single letter (i.e. "X") code. An "rt" before "Xaa<sub>1</sub>nXaa<sub>2</sub>" means "reverse transcriptase". An "s" means an envelope gene. The amino acid residues for HBV DNA polymerase are numbered with the residue methionine in the motif Tyr Met Asp Asp (YMDD) being residue number 204 (Stuyver et al., Hepatology 33: 751-757, 2001). The amino acid residues for hepatitis B virus surface antigen are number according to Norder et al. (J. Gen. Virol. 74: 341-1348, 1993). Both single and three letter abbreviations are used to define amino acid residues and these are summarized in Table 2.

In accordance with the present invention, the selection of HBV variants is identified in patients (Patient A to L) with chronic HBV infection treated with ADV. Patient E is a nonresponder to ADV. Variants of HBV are identified during ADV or combination ADV and LMV treatment with mutations in the HBV DNA polymerase gene which reduce the sensitivity of HBV to this nucleoside analog. Consequently, HBV rt variants are contemplated which are resistant to, or which exhibit reduced sensitivity to, ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and

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FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combinations thereof. Corresponding mutations in the surface antigen also occur. The identification of these HBV variants is important for the development of assays to monitor ADV, LMV, FTC and/or TFV resistance and/or resistance to other nucleoside or nucleotide analogs or other anti-HBV agents or combinations thereof and to screen for agents which are useful as alternative therapeutic agents.

Reference herein to "anti-HBV agents" includes nucleoside and nucleotide analogs as well as immunological reagents (e.g. antibodies to HBV surface components) and chemical, proteinaceous and nucleic acid agents which inhibit or otherwise interfere with viral replication, maintenance, infection, assembly or release.

The detection of such HBV variants is particularly important in the management of therapeutic protocols including the selection of appropriate agents for treating HBV infection. The method of this aspect of the present invention is predicated in part on monitoring the development in a subject of an increased HBV load in the presence of a nucleoside or nucleotide analog or other anti-HBV agents or combinations thereof. The clinician is then able to modify an existing treatment protocol or select an appropriate treatment protocol accordingly.

Accordingly, one aspect of the present invention is directed to an isolated HBV variant comprising a nucleotide mutation in a gene encoding a DNA polymerase resulting in at least one amino acid addition, substitution and/or deletion to the DNA polymerase and which exhibits decreased sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combinations thereof. The variant HBV comprises a mutation in an overlapping open reading frame in its genome in a region defined by one or more of domains F and G and domain A through to E of HBV DNA polymerase.

Another aspect of the present invention provides an isolated HBV variant comprising a nucleotide mutation in the S gene resulting in at least one amino acid addition, substitution and/or deletion to the surface antigen and which exhibits decreased sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combinations thereof.

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Useful mutants in the rt region include, in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes rtT128S rtL180M, rtM204V rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, i

Other HBV variants are also contemplated with mutations rtT38K (in the F domain of the DNA polymerase), rtR55H (located between the F and A domains), rtS/T78S, rtV80L (these are located within the A domain), rtN/S118N, rtI122V rtN/K139K, rtE142V (located between the A and B domains, rtA181V, rtA181T (these are located in the B domain),; rtI187V (located between the B and C domains), rtA/V200V (Located in the C Domain), rtV214A rtQ/P/S/Stop215S, rtQ215S, rt E/K218E (located between the C and D domains), rtn236T, rtH237H/P rtN/H238H, rtN238T, rtN238T/A (these are located in the D domain), rtY245H (located between the D and E domains), and rtV253G (located in the E Domain) or a combination thereof or an equivalent mutation.

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Useful mutations in the S gene include, in one embodiment include sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sQ101R, sI195M, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, or a combination thereof or an equivalent mutation.

The present invention further contemplates a method for determining the potential for an HBV to exhibit reduced sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof by isolating DNA or corresponding mRNA from the HBV and screening for a mutation in the nucleotide sequence encoding HBV DNA polymerase resulting in at least one amino acid substitution, deletion and/or addition in any one or more of domains F and G and domains A through to E or a region proximal thereto of the DNA polymerase and associated with resistance or decreased sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof. The presence of such a mutation is an indication of the likelihood of resistance to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and 30

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FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

The present invention also provides a composition comprising a variant HBV resistant to 5 ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV, ADV and LMV and FTC and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof or an HBV surface antigen from the variant HBV or a recombinant or derivative form thereof or its chemical equivalent and one or more pharmaceutically acceptable carriers and/or diluents.

Yet another aspect of the present invention provides a use of the aforementioned composition or a variant HBV comprising a nucleotide mutation in a gene encoding a DNA polymerase resulting in at least one amino acid addition, substitution and/or deletion to the DNA polymerase and a decreased sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof in the manufacture of a medicament for the treatment and/or prophylaxis of hepatitis B virus infection.

The present invention also contemplates a method for determining whether an HBV strain exhibits reduced sensitivity to a nucleoside or nucleotide analog or other anti-HBV agents or by isolating DNA or corresponding mRNA from the HBV and screening for a mutation in the nucleotide sequence encoding the DNA polymerase wherein the presence of the following mutations in the rt region: in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another embodiment includes rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, or combinations thereof or an equivalent one or more other mutation is indicative of a variant which exhibits a decreased sensitivity to ADV, LMV,TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

Still a further methodology comprises screening for a mutation in the nucleotide sequence encoding the envelope genes (s) wherein the presence of the following mutations in the S gene: , in one embodiment include sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, 20 sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sQ101R, sI195M, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, or combinations thereof or an equivalent one or more other mutation is indicative of a variant which exhibits a decreased sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV, and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof. 30

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Preferably, the variants are in an isolated form such that they have undergone at least one purification step away from naturally occurring body fluid. Alternatively, the variants may be maintained in isolated body fluid or may be in DNA form. The present invention also contemplates infectious molecular clones comprising the genome or parts thereof from a variant HBV. The detection of HBV or its components in cells, cell lysates, cultured supernatant fluid and bodily fluid may be by any convenient means including any nucleic acid-based detection means, for example, by nucleic acid hybridization techniques or via one or more polymerase chain reactions (PCRs). The term "bodily fluid" includes any fluid derived from the blood, lymph, tissue or organ systems including serum, whole blood, biopsy and biopsy fluid, organ explants and organ suspension such as liver suspensions.

Another aspect of the present invention is directed to a variant HBV comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or a truncation compared to a surface antigen from a reference or wild type HBV and wherein an antibody generated to the reference or wild type surface antigen exhibits an altered immunological profile relative to the HBV variant. One altered profile includes a reduced capacity for neutralizing the HBV. More particularly, the surface antigen of the variant HBV exhibits an altered immunological profile compared to a pre-treatment HBV where the variant HBV is selected for by a nucleoside or nucleotide analog or other anti-HBV agents of the HBV DNA polymerase. The variant HBV of this aspect of the invention may also comprise a nucleotide sequence comprising a single or multiple nucleotide substitution, addition and/or deletion compared to a pre-treatment HBV.

The present invention extends to an isolated HBsAg or a recombinant form thereof or derivative or chemical equivalent thereof corresponding to the variant HBV. Generally, the HBsAg or its recombinant or derivative form or its chemical equivalent comprises an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or a truncation compared to an HBsAg from a reference HBV and wherein an antibody directed to a reference HBV exhibits an altered immunological profile to an HBV

carrying said variant HBsAg. In one embodiment, the altered immunological profile comprises a reduction in the ability to neutralize the variant HBV.

Another aspect of the present invention contemplates a method for detecting an agent which exhibits inhibitory activity to an HBV by generating a genetic construct comprising a replication competent-effective amount of the genome from the HBV contained in a plasmid vector and then transfecting said cells with said construct, contacting the cells, before, during and/or after transfection, with the agent to be tested, culturing the cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release virus or virus-like particles if resistant to said agents; and the subjecting the cells, cell lysates or culture supernatant fluid to viralcomponent-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of the agent. In a preferred embodiment, the plasmid vector in a baculovirus vector and the method comprises generating a genetic construct comprising a replication competent-effective amount of the genome from the HBV contained in or fused to an amount of a baculovirus genome effective to infect cells and then infecting said cells with said construct, contacting the cells, before, during and/or after infection, with the agent to be tested, culturing the cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release virus or virus-like particles if resistant to said agent and then subjecting the cells, cell lysates or culture supernatant fluid to viral- or viral-component-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of the agent.

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In connection with these methods, the plasmid vector may include genes encoding part or all of other viral vectors such as baculovirus vectors or adenovirus vectors (see Ren and Nassal, *J. Virol.* 75(3): 1104-1116, 2001).

30 In an alternative embodiment, the method comprises generating a continuous cell line comprising an infectious copy of the genome of the HBV in a replication competent

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effective amount such that said infectious HBV genome is stably integrated into said continuous cell line such as but not limited to the 2.2.15 or AD cell line, contacting the cells with the agent to be tested, culturing the cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release virus or virus-like particles if resistant to the agent and then subjecting the cells, cell lysates or culture supernatant fluid to viral- or viral-component-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of the agent.

In an alternative embodiment, the present invention also contemplates a method for detecting an agent which exhibits inhibitory activity to an HBV polymerase in an *in vitro* polymerase assay. The HBV polymerase activity can be examined using established assays (Gaillard et al., Antimicrob Agents Chemother. 46(4): 1005-1013, 2002; Xiong et al., Hepatology. 28(6): 1669-73, 1998). The HBV polymerase may be a wild-type or reference 15 HBV polymerase or mutant HBV polymerase.

The identification of viral variants enables the production of vaccines comprising particular recombinant viral components such as polymerases or envelope genes PreS1, PreS2, S encoding for L, M, S proteins as well as therapeutic vaccines comprising defective HBV variants. Rational drug design may also be employed to identify or generate therapeutic molecules capable of interacting with a polymerase or envelope genes PreS1, PreS2, S encoding for L, M, S proteins or other component of the HBV. Such drugs may also have diagnostic potential. In addition, defective HBV variants may also be used as therapeutic compositions to generate an immune response against the same, similar or homologous viruses. Alternatively, antibodies generated to the HBV variants or surface components thereof may be used in passive immunization of subjects against infection by HBV variants or similar or homologous viruses. Furthermore, agents such as nucleoside or nucleotide analogs, RNAi or siRNA molecules, antisense or sense oligonucleotides, chemical or proteinaceous molecules having an ability to down-regulate the activity of a component of HBV and inhibit replication, maintenance, infection, assembly or release are contemplated by the present invention.

A summary of the abbreviations used throughout the subject specification are provided in Table 3.

A summary of sequence identifiers used throughout the subject specification is provided in Table 1.

TABLE 1
Summary of sequence identifiers

SEQUENCE ID NO:	DESCRIPTION	
1	Formula I	
2	Formula II	
3	OS1 primer	
4	TTA3 primer	
5	JM primer	
6	TTA4 primer	
7	OS2 primer	
8	sense primer	
9	antisense primer	
10	internal regions primer	
11	internal regions primer	
12	PC1 forward primer	
13	PC2 reverse primer	
14	HBV-specific molecular beacon primer	

TABLE 2
Single and three letter amino acid abbreviations

Amino Acid	Three-letter Abbreviation	One-letter symbol
Alanine	Ala	A
Arginine	Arg	R
Asparagine	Asn	N
Aspartic acid	Asp	D
Cysteine	Cys	С
Glutamine	Gln	Q
Glutamic acid	Glu	Е
Glycine	Gly	G
Histidine	His	Н
Isoleucine	Ile	I
Leucine	Leu	L
Lysine	Lys	K
Methionine	Met	M
Phenylalanine	Phe	F
Proline	Pro	P
Serine	Ser	S
Threonine	The	T
Tryptophan	Trp	W
Tyrosine	Tyr	Y
Valine	Val	V
Any residue	Xaa	Х

TABLE 3

Abbreviations

ABBREVIATION	DESCRIPTION	
3TC	(LMV); (-)-β-2'-deoxy-3'-thiacytidine	
ADV	adefovir dipivoxil	
DAPD	diaminopurine dioxalone	
DXG	dioxolane guanine	
ETV	entecavir	
FAM	famciclovir	
FCV	famciclovir	
FTC	emtricitabine	
HBIG	hepatitis B immunoglobulin	
HBsAg	hepatitis B surface antigen	
HBV	hepatitis B virus	
LMV	lamividuine	
PMEA	9-[phosphonyl-methoxyethyl]-adenine; adefovir	
PMPA	9-R-(2-phosphonomethoxypropyl)adenine	
RNase	ribonuclease	
rt ("rt" before "Xaa <sub>1</sub> nXaa <sub>2</sub> " means reverse transcriptase)	reverse transcriptase	
s (as used in a mutation, e.g. sF134V)	envelope gene	
TFV	tenofovir disoproxil fumarate	
YMDD	Tyr Met Asp Asp-a motif in the polymerase protein; wh the Met residue is designated residue number 204 of the reverse transcriptase	

### BRIEF DESCRIPTION OF THE FIGURES

Figure 1 is a diagrammatic representation showing the partially double stranded DNA HBV genome showing the overlapping open reading frames encoding surface (S), core (C), polymerase (P) and X gene.

Figure 2 is a diagrammatic representation of the chemical structure of ADV.

Figure 3 is a diagrammatic representation of a computer system for determining the value of a variant HBV.

Figure 4 is a representation showing comparison of the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in sequential samples from Patient A during ADV treatment.

Figure 5 is a representation showing comparison of the deduced amino acid sequence of the catalytic region of the polymerase gene in sequential samples from Patient A during ADV therapy.

Figure 6 is a representation showing comparison of the deduced amino acid sequence of the envelope gene in sequential samples from Patient A during ADV therapy.

Figure 7 is a representation showing comparison of the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in sequential samples from Patient B during ADV and LMV treatment.

Figure 8 is a representation showing comparison of the deduced amino acid sequence of the catalytic region of the polymerase gene in sequential samples from Patient B during ADV and LMV therapy.

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- Figure 9 is a representation showing comparison of the deduced amino acid sequence of the envelope gene in sequential samples from Patient B during ADV and LMV therapy.
- Figure 10 is a representation the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient C during ADV treatment.
  - Figure 11 is a representation the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient C during ADV therapy.
- 10 Figure 12 is a representation the deduced amino acid sequence of the envelope gene in samples from Patient C during ADV therapy.
  - Figure 13 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient D during ADV treatment.
  - Figure 14 is a representation the deduced amino acid sequence of the catalytic region of the polymerase gene in sequential samples from Patient D during ADV therapy.
- Figure 15 is a representation showing comparison of the deduced amino acid sequence of the envelope gene in sequential samples from Patient D during ADV therapy.
  - Figure 16 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient E during ADV treatment.
- Figure 17 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient E during ADV therapy.
  - Figure 18 is a representation showing the deduced amino acid sequence of the envelope gene in samples from Patient E during ADV therapy.

- Figure 19 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient F during ADV treatment.
- Figure 20 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient F during ADV therapy.
  - Figure 21 is a representation showing the deduced amino acid sequence of the envelope gene in samples from Patient F during ADV therapy.
- 10 Figure 22 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient G during ADV treatment.
  - Figure 23 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient G during ADV therapy.
  - Figure 24 is a representation showing the deduced amino acid sequence of the envelope gene in samples from Patient G during ADV therapy.
- Figure 25 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient H during ADV treatment.
  - Figure 26 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient H during ADV therapy.
- 25 Figure 27 is a representation showing the deduced amino acid sequence of the envelope gene in samples from Patient H during ADV therapy.
  - Figure 28 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient I during ADV treatment.

- Figure 29 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient I during ADV therapy.
- Figure 30 is a representation showing the deduced amino acid sequence of the envelope gene in samples from Patient I during ADV therapy.
  - Figure 31 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient J during ADV treatment.
- Figure 32 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient J during ADV therapy.
  - Figure 33 is a representation showing the deduced amino acid sequence of the envelope gene in samples from Patient J during ADV therapy.
  - Figure 34 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient K during ADV treatment.
- Figure 35 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient K during ADV therapy.
  - Figure 36 is a representation showing the deduced amino acid sequence of the envelope gene in samples from Patient K during ADV therapy.
- Figure 37 is a representation showing the HBV nucleotide sequence encoding the catalytic region of the polymerase gene in samples from Patient L during ADV treatment.
  - Figure 38 is a representation showing the deduced amino acid sequence of the catalytic region of the polymerase gene in samples from Patient L during ADV therapy.

Figure 39 is a representation showing the deduced amino acid sequence of the envelope . gene in samples from Patient L during ADV therapy..

### DETAILED DESCRIPTION OF THE INVENTION

The present invention is predicated in part on the identification and isolation of nucleoside or nucleotide analog-resistant variants of HBV following treatment of patients with either 5 ADV or LMV or more particularly ADV and LMV, or optionally other nucleoside analogs or nucleotide analogs or other anti-HBV agents such as TFV or FTC. In particular, ADV or ADV and LMV treated patients gave rise to variants of HBV exhibiting decreased or reduced sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV. Reference herein to "decreased" or "reduced" in relation to sensitivity to ADV and/or LMV and/or FTC and/or TFV includes and encompasses a complete or substantial resistance to the nucleoside or nucleotide analog or other anti-HBV agents as well as partial resistance and includes a replication rate or replication efficiency which is more than a wild-type in the presence of a nucleoside or nucleotide analog or other anti-HBV agents. In one aspect, this is conveniently measured by an increase in viral load during treatment, or alternatively, there is no substantial decrease in HBV DNA viral load from pre-treatment HBV DNA levels during treatment (i.e., non-response to treatment).

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Before describing the present invention in detail, it is to be understood that unless otherwise indicated, the subject invention is not limited to specific formulations of components, manufacturing methods, dosage regimens, or the like, as such may vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only and is not intended to be limiting.

It must be noted that, as used in the subject specification, the singular forms "a", "an" and "the" include plural aspects unless the context clearly dictates otherwise. Thus, for example, reference to "a nucleoside or nucleotide analog" includes a single analog, as well as two or more analogs; reference to "an HBV variant" includes reference to two or more HBV variants; and so forth.

In describing and claiming the present invention, the following terminology is used in accordance with the definitions set forth below.

The terms "analog", "compound", "active agent", "pharmacologically active agent", "medicament", "active" and "drug" are used interchangeably herein to refer to a chemical compound that induces a desired effect such as inhibit viral replication, infection, maintenance, assembly and/or the function of an enzyme such as HBV DNA polymerase. The terms also encompass pharmaceutically acceptable and pharmacologically active ingredients of those active agents specifically mentioned herein including but not limited to salts, esters, amides, prodrugs, active metabolites, analogs and the like. When the terms "analog", "compound", "active agent", "pharmacologically active agent", "medicament", "active" and "drug" are used, then it is to be understood that this includes the active agent per se as well as pharmaceutically acceptable, pharmacologically active salts, esters, amides, prodrugs, metabolites, analogs, etc.

The present invention contemplates, therefore, compounds useful in inhibiting HBV replication, infection, maintenance, assembly and/or the function of an enzyme such as HBV DNA polymerase. Reference to an "analog", "compound", "active agent", "pharmacologically active agent", "medicament", "active" and "drug" such as ADV, LMV, FTC and/or TFV includes combinations of two or more actives such as ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV. A "combination" also includes a two-part or more such as a multi-part anti-HBV therapeutic composition where the agents are provided separately and given or dispensed separately or admixed together prior to dispensation.

The terms "effective amount" and "therapeutically effective amount" of an agent as used herein mean a sufficient amount of the agent to provide the desired therapeutic or physiological effect of inhibiting HBV replication, infection, maintenance, assembly

and/or the function of an enzyme such as HBV DNA polymerase. Furthermore, an "effective HBV-inhibiting amount" or "effective symptom-ameloriating amount" of an agent is a sufficient amount of the agent to directly or indirectly inhibit replication, infection, maintenance, assembly and/or the function of an enzyme such as HBV DNA polymerase. Undesirable effects, e.g. side effects, are sometimes manifested along with the desired therapeutic effect; hence, a practitioner balances the potential benefits against the potential risks in determining what is an appropriate "effective amount". The exact amount required will vary from subject to subject, depending on the species, age and general condition of the subject, mode of administration and the like. Thus, it may not be possible to specify an exact "effective amount". However, an appropriate "effective amount" in any individual case may be determined by one of ordinary skill in the art using only routine experimentation.

By "pharmaceutically acceptable" carrier, excipient or diluent is meant a pharmaceutical vehicle comprised of a material that is not biologically or otherwise undesirable, i.e. the material may be administered to a subject along with the selected active agent without causing any or a substantial adverse reaction. Carriers may include excipients and other additives such as diluents, detergents, coloring agents, wetting or emusifying agents, pH buffering agents, preservatives, and the like.

Similarly, a "pharmacologically acceptable" salt, ester, emide, prodrug or derivative of a compound as provided herein is a salt, ester, amide, prodrug or derivative that this not

biologically or otherwise undesirable.

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The terms "treating" and "treatment" as used herein refer to reduction in severity and/or frequency of symptoms, elimination of symptoms and/or underlying cause, prevention of the occurrence of symptoms and/or their underlying cause, and improvement or remediation of damage in relation to HBV infection. Thus, for example, "treating" a patient involves prevention of HBV infection as well as treatment of a clinically HBV symptomatic individual by inhibiting HBV replication, infection, maintenance, assembly and/or the function of an enzyme such as HBV DNA polymerase. Thus, for example, the

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present method of "treating" a patient with HBV infection or with a propensity for one to develop encompasses both prevention of HBV infection as well as treating HBV infection or symptoms thereof. In any event, the present invention contemplates the treatment or prophylaxis of HBV infection.

"Patient" as used herein refers to an animal, preferably a mammal and more preferably a primate including a lower primate and even more preferably, a human who can benefit from the formulations and methods of the present invention. A patient regardless of whether a human or non-human animal may be referred to as an individual, subject, animal, host or recipient. The compounds and methods of the present invention have applications in human medicine, veterinary medicine as well as in general, domestic or wild animal husbandry. For convenience, an "animal" includes an avian species such as a

poultry bird (including ducks, chicken, turkeys and geese), an aviary bird or game bird. The condition in a non-human animal may not be a naturally occurring HBV infection but

15 HBV-like infection may be induced.

As indicated above, the preferred animals are humans, non-human primates such as marmossets, baboons, orangatangs, lower primates such as tupia, livestock animals, laboratory test animals, companion animals or captive wild animals. A human is the most preferred target. However, non-human animal models may be used.

Examples of laboratory test animals include mice, rats, rabbits, guinea pigs and hamsters. Rabbits and rodent animals, such as rats and mice, provide a convenient test system or animal model as do primates and lower primates. Livestock animals include sheep, cows, pigs, goats, horses and donkeys. Non-mammalian animals such as avian species, zebrafish, amphibians (including cane toads) and *Drosophila* species such as *Drosophila* melanogaster are also contemplated. Instead of a live animal model, a test system may also comprise a tissue culture system.

Accordingly, one aspect of the present invention is directed to an isolated Hepatitis B virus (HBV) variant wherein said variant comprises a nucleotide mutation in a gene encoding a

DNA polymerase resulting in at least one amino acid addition, substitution and/or deletion to said DNA polymerase and wherein said variant exhibits decreased sensitivity to one of ADV, LMV,TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV or optionally other nucleoside analogs or other anti-HBV agents or a combination thereof.

An "anti-HBV agent" includes a nucleoside or nucleotide analog, protein, chemical compound, RNA or DNA or RNAi or siRNA oligonucleotide.

Preferably, the decreased sensitivity is in respect of ADV. Alternatively, the decreased sensitivity is in respect of LMV. Alternatively, the decreased sensitivity is in respect of TFV. Alternatively, the decreased sensitivity is in respect of FTC. Alternatively, the decreased sensitivity is in respect of ADV and LMV. Alternatively, the decreased sensitivity is in respect of ADV and TFV. Alternatively, the decreased sensitivity is in respect of ADV and FTC. Alternatively, the decreased sensitivity is in respect to FTC and TFV. Alternatively, the decreased sensitivity is in respect to FTC and TFV. Alternatively, the decreased sensitivity is in respect of ADV and LMV. Alternatively, the decreased sensitivity is in respect to ADV and TFV and FTC. Alternatively, the decreased sensitivity is in respect to LMV and TFV and FTC. Alternatively, the decreased sensitivity is in respect to LMV and TFV. Alternatively, the decreased sensitivity is in respect of ADV and LMV and FTC. Alternatively, the decrease sensitivity is in respect of ADV and LMV and FTC. Alternatively, the decrease sensitivity is in respect of ADV and TFV and FTC. Alternatively, the decrease sensitivity is in respect of ADV and FTC and TFV and LMV.

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Reference herein to "anti-HBV agents" includes nucleoside and nucleotide analogs as well as immunological reagents (e.g. antibodies to HBV surface components) and chemical, proteinaceous and nucleic acid agents which inhibit or otherwise interfere with viral replication, maintenance, infection, assembly or release. Reference herein to "nucleic acid" includes reference to a sense or antisense molecule, RNA or DNA, oligonucleotides and RNAi and siRNA molecules and complexes containing same.

In addition to a mutation in the gene encoding DNA polymerase, due to the overlapping nature of the HBV genome (Figure 1), a corresponding mutation may also occur in the gene encoding the S gene encoding the surface antigen (HBsAg) resulting in reduced interactivity of immunological reagents such as antibodies and immune cells to HBsAg. The reduction in the interactivity of immunological reagents to a viral surface component generally includes the absence of immunological memory to recognize or substantially recognize the viral surface component. The present invention extends, therefore, to an HBV variant exhibiting decreased sensitivity to ADV, LMV, TFV or FTC; ADV and 10 LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV or a reduced interactivity of an immunological reagent to HBsAg wherein the variant is selected for following ADV and/or LMV combination or sequential treatment. The term "sequential" in this respect means ADV followed by LMV and/or TFV, and /or FTC, LMV followed by ADV and/or TFV, and /or FTC, or multiple sequential administrations of each of ADV, LMV and/or TFV, and /or FTC.

A viral variant may, therefore, carry mutation only in the DNA polymerase gene or both in the DNA polymerase gene and the S gene. The term "mutation" is to be read in its broadest context and includes multiple mutations.

The present invention extends to a mutation and any domain of the HBV DNA polymerase and in particular regions F and G, and domains A through to E provided said mutation leads to decreased sensitivity to ADV and/ or LMV and/or TFV or combinations thereof. Regions F and G of the HBV DNA polymerase is defined by the amino acid sequence set forth in Formula I below [SEQ ID NO:1]:

#### **FORMULA I**

L, B<sub>1</sub>, B<sub>2</sub>, D, W, G, P, C, B<sub>3</sub>, B<sub>4</sub>, H, G, B<sub>5</sub>, H, B<sub>6</sub>, I, R, B<sub>7</sub>, P, R, T, P, B<sub>8</sub>, R, V, B<sub>9</sub>, G, G, V, F, L, V, D, K, N, P, H, N, T, B<sub>10</sub>, E, S, B<sub>11</sub>, L, B<sub>12</sub>, V, D, F, S, Q, F, S, R, G, B<sub>13</sub>, B<sub>14</sub>, B 15, V, S, W, P, K, F, A, V, P, N, L, B<sub>16</sub>, S, L, T, N, L, L, Sx

### wherein:

**B**<sub>16</sub>

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is L, or R, or I  $\mathbf{B_1}$ is E, or D 10. B<sub>2</sub> is T, or D, or A, or N, or Y  $B_3$ is E, or D  $B_4$ is E, or K, or Q B<sub>5</sub> is H, or R, or N,  $B_6$ is I, or T 15  $B_7$ is A, or S  $\mathbf{B_8}$ is T or R  $\mathbf{B}_{9}$ is A, or T, or S  $B_{10}$ is R, or T  $B_{11}$ is V, or G B<sub>12</sub> 20 is S, or I, or T, or N, or V B<sub>13</sub> is T, or S, or H, or Y B<sub>14</sub> is R, or H, or K, or Q B<sub>15</sub> is Q, or P;

and wherein Sx is designated as amino acid 74.

In this specification, reference is particularly made to the conserved regions of the DNA polymerase as defined by domains A to E. Regions A to E are defined by the amino acid sequence set forth in Formula II below [SEQ ID NO:2] (and in Australian Patent No. 734831):

### **FORMULA II**

S Z<sub>1</sub> L S W L S L D V S A A F Y H Z<sub>2</sub> P L H P A A M P H L L Z<sub>3</sub> G S S G L Z<sub>4</sub> R Y V A

5 R L S S Z<sub>5</sub> S Z<sub>6</sub> Z<sub>7</sub> X N Z<sub>8</sub> Q Z<sub>9</sub> Z<sub>10</sub> X X X Z<sub>11</sub> L H Z<sub>12</sub> Z<sub>13</sub> C S R Z<sub>14</sub> L Y V S L Z<sub>15</sub> L L Y

Z<sub>16</sub> T Z<sub>17</sub> G Z<sub>18</sub> K L H L Z<sub>19</sub> Z<sub>20</sub> H P I Z<sub>21</sub> L G F R K Z<sub>22</sub> P M G Z<sub>23</sub> G L S P F L L A Q F

T S A I Z<sub>24</sub> Z<sub>25</sub> Z<sub>26</sub> Z<sub>27</sub> Z<sub>28</sub> R A F Z<sub>29</sub> H C Z<sub>30</sub> Z<sub>31</sub> F Z<sub>32</sub> Y M<sup>x</sup> D D Z<sub>33</sub> V L G A Z<sub>34</sub> Z<sub>35</sub> Z<sub>36</sub>

Z<sub>37</sub> H Z<sub>38</sub> E Z<sub>39</sub> L Z<sub>40</sub> Z<sub>41</sub> Z<sub>42</sub> Z<sub>43</sub> Z<sub>44</sub> Z<sub>45</sub> Z<sub>46</sub> L L Z<sub>47</sub> Z<sub>48</sub> G I H L N P Z<sub>49</sub> K T K R W G Y

S L N F M G Y Z<sub>50</sub> I G

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#### wherein:

- X is any amino acid;
- $Z_1$  is N or D;
- 15  $\mathbb{Z}_2$  is I or P;
  - $Z_3$  is I or V;
  - $Z_4$  is S or D;
  - $Z_5$  is T or N;
  - Z<sub>6</sub> is R or N;
- 20 Z<sub>7</sub> is N or I;
  - $Z_8$  is N or Y or H;
  - Z<sub>9</sub> is H or Y;
  - $Z_{10}$  is G or R;
  - $Z_{11}$  is D or N;
- 25  $Z_{12}$  is D or N;
  - $Z_{13}$  is S or Y;
  - $Z_{14}$  is N or Q;
  - $Z_{15}$  is L or M;
  - $Z_{16}$  is K or Q;
- 30  $Z_{17}$  is Y or F;
  - $Z_{18}$  is R or W;

 $Z_{19}$ is Y or L; is S or A;  $Z_{20}$ is I or V;  $Z_{21}$ is I or L;  $Z_{22}$  $Z_{23}$ is V or G; 5 is C or L;  $Z_{24}$ is A or S;  $Z_{25}$ is V or M;  $Z_{26}$ is V or T;  $Z_{27}$ is R or C; 10  $Z_{28}$ is F or P;  $Z_{29}$  $Z_{30}$ is L or V;  $Z_{31}$ is A or V;  $Z_{32}$ is S or A; 15 Z<sub>33</sub> is V or L or M; is K or R;  $Z_{34}$ is S or T;  $Z_{35}$ is V or G;  $Z_{36}$ is Q or E;  $Z_{37}$ is L or S or R; 20  $\mathbb{Z}_{38}$ is S or F;  $Z_{39}$ is F or Y;  $Z_{40}$ is T or A;  $Z_{41}$ is A or S;  $Z_{42}$ is V or I;  $Z_{43}$ 25 **Z**44 is T or C; Z45 is N or S; is F or V; Z46 Z47 is S or D; is L or V;  $\mathbb{Z}_{48}$ 30

is N or Q;

Z49

Z<sub>50</sub> is V or I; and

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M<sup>x</sup> is amino acid 204;

and wherein the first S is designated as amino acid 75.

Preferably, the mutation results in an altered amino acid sequence in any one or more of domains F and G, and domains A through to E or regions proximal thereto of the HBV DNA polymerase.

Another aspect of the present invention provides an HBV variant comprising a mutation in an overlapping open reading frame in its genome wherein said mutation is in a region defined by one or more of domains F and G, and domains A through to E of HBV DNA polymerase and wherein said variant exhibits decreased sensitivity to ADV, LMV, TFV or FTC; ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV;

15 FTC and LMV; ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; and/or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents.

In a related embodiment, there is provided an HBV variant comprising a mutation in the nucleotide sequence encoding a DNA polymerase resulting in an amino acid addition, substitution and/or deletion in said DNA polymerase in one or more amino acids as set

forth in Formula I [SEQ ID NO:1] and/or Formula II [SEQ ID NO:2]:

#### **FORMULA I**

25 L, B<sub>1</sub>, B<sub>2</sub>, D, W, G, P, C, B<sub>3</sub>, B<sub>4</sub>, H, G, B<sub>5</sub>, H, B<sub>6</sub>, I, R, B<sub>7</sub>, P, R, T, P, B<sub>8</sub>, R, V, B<sub>9</sub>, G, G, V, F, L, V, D, K, N, P, H, N, T, B<sub>10</sub>, E, S, B<sub>11</sub>, L, B<sub>12</sub>, V, D, F, S, Q, F, S, R, G, B<sub>13</sub>, B<sub>14</sub>, B<sub>15</sub>, V, S, W, P, K, F, A, V, P, N, L, B<sub>16</sub>, S, L, T, N, L, L, Sx

wherein:

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B<sub>1</sub> is L, or R, or I

 $B_2$  is E, or D

B<sub>3</sub> is T, or D, or A, or N, or Y

B<sub>4</sub> is E, or D

B<sub>5</sub> is E, or K, or Q

5  $B_6$  is H, or R, or N,

B<sub>7</sub> is I, or T

B<sub>8</sub> is A, or S

 $B_9$  is T or R

 $B_{10}$  is A, or T, or S

10 B<sub>11</sub> is R, or T

 $B_{12}$  is V, or G

 $B_{13}$  is S, or I, or T, or N, or V

 $B_{14}$  is T, or S, or H, or Y

 $B_{15}$  is R, or H, or K, or Q

15  $B_{16}$  is Q, or P;

and

20 FORMULA II

S Z<sub>1</sub> L S W L S L D V S A A F Y H Z<sub>2</sub> P L H P A A M P H L L Z<sub>3</sub> G S S G L Z<sub>4</sub> R Y V A R L S S Z<sub>5</sub> S Z<sub>6</sub> Z<sub>7</sub> X N Z<sub>8</sub> Q Z<sub>9</sub> Z<sub>10</sub> X X X Z<sub>11</sub> L H Z<sub>12</sub> Z<sub>13</sub> C S R Z<sub>14</sub> L Y V S L Z<sub>15</sub> L L Y Z<sub>16</sub> T Z<sub>17</sub> G Z<sub>18</sub> K L H L Z<sub>19</sub> Z<sub>20</sub> H P I Z<sub>21</sub> L G F R K Z<sub>22</sub> P M G Z<sub>23</sub> G L S P F L L A Q F T S A I Z<sub>24</sub> Z<sub>25</sub> Z<sub>26</sub> Z<sub>27</sub> Z<sub>28</sub> R A F Z<sub>29</sub> H C Z<sub>30</sub> Z<sub>31</sub> F Z<sub>32</sub> Y M<sup>x</sup> D D Z<sub>33</sub> V L G A Z<sub>34</sub> Z<sub>35</sub> Z<sub>36</sub> Z<sub>37</sub> H Z<sub>38</sub> E Z<sub>39</sub> L Z<sub>40</sub> Z<sub>41</sub> Z<sub>42</sub> Z<sub>43</sub> Z<sub>44</sub> Z<sub>45</sub> Z<sub>46</sub> L L Z<sub>47</sub> Z<sub>48</sub> G I H L N P Z<sub>49</sub> K T K R W G Y S L N F M G Y Z<sub>50</sub> I G

wherein:

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X is any amino acid;

is N or D;

 $Z_{1}$ 

 $\mathbb{Z}_2$ is I or P;  $Z_3$ is I or V;  $Z_4$ is S or D; 5  $Z_5$ is T or N;  $Z_6$ is R or N;  $Z_7$ is N or I;  $Z_8$ is N or Y or H; Z<sub>9</sub> is H or Y; 10  $Z_{10}$ is G or R;  $Z_{11}$ is D or N;  $Z_{12}$ is D or N;  $Z_{13}$ is S or Y;  $Z_{14}$ is N or Q; 15 Z<sub>15</sub> is L or M;  $Z_{16}$ is K or Q;  $Z_{17}$ is Y or F;  $Z_{18}$ is R or W;  $Z_{19}$ is Y or L; 20  $Z_{20}$ is S or A;  $Z_{21}$ is I or V;  $Z_{22}$ is I or L;  $\mathbb{Z}_{23}$ is V or G;  $Z_{24}$ is C or L;  $Z_{25}$ is A or S; 25  $Z_{26}$ is V or M;  $Z_{27}$ is V or T; is R or C;  $Z_{28}$  $Z_{29}$ is F or P;

is L or V;

is A or V;

 $Z_{30}$ 

 $Z_{31}$ 

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Z_{32}
               is S or A;
      Z_{33}
               is V or L or M;
               is K or R;
      Z_{34}
      Z_{35}
               is S or T;
      Z_{36}
               is V or G;
      Z_{37}
               is Q or E;
               is L or S or R;
      Z_{38}
               is S or F;
      Z_{39}
               is F or Y;
      Z_{40}
10
      Z_{41}
               is T or A;
      Z_{42}
               is A or S;
      Z_{43}
               is V or I;
      Z_{44}
               is T or C;
               is N or S;
      Z_{45}
15
      Z_{46}
               is F or V;
      Z47
               is S or D;
               is L or V;
      Z_{48}
      Z_{49}
               is N or Q;
               is V or I; and
      Z_{50}
      M^{x}
               is amino acid 204;
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and wherein Sx in Formula I is designated as amino acid 74 and the first S in Formula II is designated as amino acid 75;

and wherein said variant exhibits decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; ADV and LMV and TFV; ADV and FTC and LMV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

Preferably, the decreased sensitivity is to ADV or to both ADV and LMV or to one or both of ADV and/or LMV and/or TFV and/or FTC.

Another preferred aspect of the present invention contemplates an HBV variant comprising a mutation in the nucleotide sequence encoding HBsAg resulting in an amino acid addition, substitution and/or deletion in said HBsAg in a region corresponding to the amino acid sequence set forth in Formulae I and II wherein said variant exhibits decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

More particularly, the present invention provides a variant HBV comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or a truncation compared to a surface antigen from a reference or wild-type HBV and wherein an antibody generated to the reference or wild-type surface antigen exhibits reduced capacity for neutralizing said HBV variant, said variant selected by exposure of a subject to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

The term "combination therapy" means that both combinations of ADV, LMV, FTC and/or TFV are co-administered in the same composition or simultaneously in separate compositions. The term "sequential therapy" means that the two agents are administered within seconds, minutes, hours, days or weeks of each other and in either order. Sequential therapy also encompasses completing a therapeutic course with one or other of ADV, LMV, FTC or TFV and then completing a second or third or subsequent therapeutic courses with the other of ADV, LMV, FTC or TFV.

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Accordingly, another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to LMV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Yet another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to FTC therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Still another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Even yet another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV and LMV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

10 Even still another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV and TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

A further aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to LMV and TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

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Another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject

to ADV and FTC therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Yet another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to TFV and FTC therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Still another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to FTC and LMV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

20 Even yet another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV, LMV and TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Even still another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile

compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV, LMV and TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

A further aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV, LMV and FTC therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to FTC, LMV and TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

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Yet another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV, FTC and TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

Still yet another aspect of the present invention contemplates an HBV variant comprising a surface antigen having an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or truncation compared to the pretreatment HBV and

wherein the surface antigen of the variant HBV exhibits an altered immunological profile compared to the pretreatment HBV where the said variant HBV is selected for by exposure of a subject to ADV, LMV, FTC and TFV therapy or therapy by one or more other nucleoside or nucleotide analogs or other anti-HBV agents.

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Preferably, the variants are in isolated form such that they have undergone at least one purification step away from naturally occurring body fluid. Alternatively, the variants may be maintained in isolated body fluid or may be in DNA form. The present invention also contemplates infectious molecular clones comprising the genome or parts thereof from a variant HBV. Furthermore, the present invention provides isolated components from the variant HBVs such as but not limited to an isolated HBsAg. Accordingly, the present invention provides an isolated HBsAg or a recombinant form thereof or derivative or chemical equivalent thereof, said HBsAg being from a variant HBV selected by exposure of a subject to one or more of ADV, LMV, FTC and/or TFV or optionally one or more nucleoside or nucleotide analogs or other anti-HBV agents.

More particularly, yet another aspect of the present invention is directed to an isolated variant HBsAg or a recombinant or derivative form thereof or a chemical equivalent thereof wherein said HBsAg or its recombinant or derivative form or its chemical equivalent exhibits an altered immunological profile compared to an HBsAg from a reference HBV, said HBsAg being from a variant HBV selected by exposure of a subject to one or more of ADV, LMV, FTC and/or TFV or optionally one or more nucleoside or nucleotide analogs or other anti-HBV agents.

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Even more particularly, the present invention provides an isolated variant HBsAg or a recombinant or derivative form thereof or a chemical equivalent thereof wherein said HBsAg or its recombinant or derivative form or its chemical equivalent comprises an amino acid sequence with a single or multiple amino acid substitution, addition and/or deletion or a truncation compared to an HBsAg from a reference HBV and wherein a neutralizing antibody directed to a reference HBV exhibits no or reduced neutralising activity to an HBV carrying said variant HBsAg, said HBsAg being from a variant HBV

selected by exposure of a subject to one or more of ADV, LMV, FTC and/or TFV or optionally one or more nucleoside or nucleotide analogs or other anti-HBV agents.

Preferred mutations in the HBV DNA polymerase include variants selected from patients with HBV recurrence following ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV treatment. Nucleoside or nucleotide analog or other anti-HBV agents treatment may occur in relation to a transplantation procedure (e.g. bone marrow transplantation (BMT) or OLT) or following treatment of patients diagnosed with hepatitis. Following selection of variants, viral loads are obtainable at levels similar to pretreatment levels or are increasing while on therapy.

Useful mutants in the rt region include, in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes rtT128S rtL180M, rtM204V rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, or a combination thereof or an equivalent mutation

Such HBV variants are proposed to exhibit a decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof. It should be noted that the nomenclature system for amino acid positions is based on the methionine residues in the YMDD motif being designated codon rtM204. This

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numbering system is different to that in Australian Patent No. 734831 where the methionine residue in the YMDD motif within the polymerase gene is designated codon 550. In this regard, rtL180M and rtM204V correspond to L526M and M550V, respectively, in Australian Patent No. 734831. Corresponding mutations may also occur in envelope genes such as in one or more of PreS1, PreS2 and S.

Another potential mode of action of ADV and other acyclic nucleoside phosphonates is that of immune stimulation (Calio et al., Antiviral Res. 23: 77-89, 1994). A number of mutations resulted in changes in the envelope gene detected in HBV variants which may be associated with immune escape. These changes include in one embodiment include sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sL/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, or a combination thereof or an equivalent mutation.

The identification of the variants of the present invention permits the generation of a range of assays to detect such variants. The detection of such variants may be important in identifying resistant variants to determine the appropriate form of chemotherapy and/or to monitor vaccination protocols, or develop new or modified vaccine preparations.

Still another aspect of the present invention contemplates a method for determining the potential for an HBV to exhibit reduced sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other

nucleoside or nucleotide analogs or other anti-HBV agents, said method comprising isolating DNA or corresponding mRNA from said HBV and screening for a mutation in the nucleotide sequence encoding HBV DNA polymerase resulting in at least one amino acid substitution, deletion and/or addition in any one or more of domains F and G, and A domains through to E or a region proximal thereto of said DNA polymerase and associated with resistance or decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents wherein the presence of such a mutation is an indication of the likelihood of resistance to said ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents. 15

Preferably, the assay detects one or more of the following mutations: in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another embodiment includes rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V; or combinations thereof or an equivalent one or more other mutation is indicative of a variant wherein said variant exhibits a decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and

FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

Accordingly, another aspect of the present invention produces a method for determining whether an HBV strain exhibits reduced sensitivity to a nucleoside or nucleotide analog or other anti-HBV agents, said method comprising isolating DNA or corresponding mRNA from said HBV and screening for a mutation in the nucleotide sequence encoding the DNA polymerase and/or a corresponding region of the S gene, wherein the presence of a mutation selected from, in one embodiment include sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sQ101R, sI195M, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, and in yet another embodiment include sT47A and sW172stop, in even still another embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtI204M, rtN/K139K, rtE142V, rtA/T181A rtN/S118N. rtV80L, 20 rtS/T78S. rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another embodiment includes 25 rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, or combinations thereof or an equivalent one or more other mutation is indicative of a variant which exhibits a decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC 30 and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

A further aspect of the present invention produces a method for determining whether an HBV strain exhibits reduced sensitivity to a nucleoside or nucleotide analog or other anti-HBV agents, said method comprising isolating DNA or corresponding mRNA from said HBV and screening for a mutation in the nucleotide sequence encoding the DNA polymerase and/or a corresponding region of the S gene, wherein the presence of a mutation selected from, in one embodiment, sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sQ101R, sI195M, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, and in yet another embodiment include sT47A and sW172stop, in even still another embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another embodiment includes rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, combinations thereof or an equivalent one or more other mutation is indicative of a variant which exhibits a decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

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The detection of HBV or its components in cells, cell lysates, cultured supernatant fluid and bodily fluid may be by any convenient means including any nucleic acid-based detection means, for example, by nucleic acid hybridization techniques or via one or more polymerase chain reactions (PCRs). The term "bodily fluid" includes any fluid derived from the blood, lymph, tissue or organ systems including serum, whole blood, biopsy and biopsy fluid, organ explants and organ suspension such as liver suspensions. The invention further encompasses the use of different assay formats of said nucleic acid-based detection means, including restriction fragment length polymorphism (RFLP), amplified fragment length polymorphism (AFLP), single-strand chain polymorphism (SSCP), amplification and mismatch detection (AMD), interspersed repetitive sequence polymerase chain reaction (IRS-PCR), inverse polymerase chain reaction (iPCR) and reverse transcription polymerase chain reaction (RT-PCR), amongst others. Other forms of detection include Northern blots, Southern blots, PCR sequencing, antibody procedures such as ELISA, Western blot and immunohistochemistry. A particularly useful assay includes the reagents and components required for immobilized oligonucleotide- or oligopeptide-mediated detection systems.

One particularly useful nucleic acid detection system is the reverse hybridization technique. In this technique, DNA from an HBV sample is amplified using a biotin or other ligand-labeled primer to generate a labeled amplificon. Oligonucleotides immobilized to a solid support such as a nitrocellulose film are then used to capture amplified DNA by hybridization. Specific nucleic acid fragments are identified via biotin or the ligand. Generally, the labeled primer is specific for a particular nucleotide variation to be detected. Amplification occurs only if the variation to be detected is present. There are many forms of the reverse hybridization assay and all are encompassed by the present invention.

Detecting HBV replication in cell culture is particularly useful.

This and other aspects of the present invention is particularly amenable to microarray analysis such as to identify oligonucleotides including sense and antisense molecules, RNAi or siRNA molecules or DNA or RNA-binding molecules which down-regulate genomic sequences or transcripts of HBV. Microarray analysis may also be used to identify particular mutations in the HBV genome such as within the HBV DNA polymerase-coding region or the HBsAg-coding region.

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Another aspect of the present invention contemplates a method for detecting an agent which exhibits inhibitory activity to an HBV by:

generating a genetic construct comprising a replication competent-effective
amount of the genome from the HBV contained in a plasmid vector and then transfecting
said cells with said construct;

contacting the cells, before, during and/or after transfection, with the agent to be tested;

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culturing the cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release virus or virus-like particles if resistant to said agents; and

then subjecting the cells, cell lysates or culture supernatant fluid to viral- or viral-component-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of the agent.

In a preferred embodiment, the plasmid vector may include genes encoding part or all of other viral vectors such as baculovirus or adenovirus (Ren and Nassal, 2001, *supra*) and the method comprises:

generating a genetic construct comprising a replication competent-effective amount of the genome from the HBV contained in or fused to an amount of a baculovirus genome or adenovirus genome effective to infect cells and then infecting said cells with said construct;

10 contacting the cells, before, during and/or after infection, with the agent to be tested;

culturing the cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release virus or virus-like particles if resistant to said agent; and

then subjecting the cells, cell lysates or culture supernatant fluid to viral- or viral-component-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of the agent.

In an alternative embodiment, the method comprises:

generating a continuous cell line comprising an infectious copy of the genome of the HBV in a replication competent effective amount such that said infectious HBV genome is stably integrated into said continuous cell line such as but not limited to 2.2.15 or AD;

contacting the cells with the agent to be tested;

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culturing the cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release virus or virus-like particles if resistant to the agent; and

then subjecting the cells, cell lysates or culture supernatant fluid to viral- or viral-component-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of the agent.

The above-mentioned methods are particularly useful in identifying or developing agents 10 against HBV variants such as those carrying mutations, in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another embodiment includes rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, or a combination thereof or an 20 equivalent mutation; in a further embodiment, sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 25 T6S, sT47A, sP62L, sL/F192F, sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sQ101R, sI195M, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, or a combination thereof or an equivalent mutation. 30

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Accordingly, another aspect of the present invention contemplates a method for determining whether an HBV strain exhibits reduced sensitivity to a nucleoside or nucleotide analog or other potential anti-HBV agent, said method comprising isolating DNA or corresponding mRNA from said HBV and screening for a mutation in the nucleotide sequence of the envelope genes or DNA polymerase gene selected from, in one embodiment, in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet 10 another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another embodiment includes rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, or a combination thereof or an equivalent mutation; in a further embodiment, sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sQ101R, sI195M, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, or combinations thereof or an equivalent one or more other mutation is indicative of a variant wherein said variant exhibits a decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof.

The detection of amino acid variants of DNA polymerase is conveniently accomplished by reference to the amino acid sequence shown in Formulae I and II. The polymorphisms shown represent the variations shown in various databases for active pathogenic HBV strains. Where an HBV variant comprises an amino acid different to what is represented, then such an isolate is considered a putative HBV variant having an altered DNA polymerase activity.

The present invention further contemplates agents which inhibit ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; 10 FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV resistant HBV variants. Such agents are particularly useful if long term treatment by ADV, LMV, FTC and/or TFV and/or optionally other nucleoside analogs or nucleotide analogs such as TFV is contemplated by the clinician. The agents may be DNA or RNA or proteinaceous or non-proteinaceous chemical molecules. Natural product screening such as from plants, coral and microorganisms is also contemplated as a useful potential source of masking agents as is the screening of combinatorial or chemical libraries. The agents may be in isolated form or in the form of a pharmaceutical composition or formulation and may be administered in place of or sequentially or simultaneously with a nucleoside or nucleotide analog. Furthermore, rationale drug design is contemplated including solving the crystal or NMR structure of, for example, HBV DNA polymerase and designing agents which can bind to the enzyme's active site. This approach may also be adapted to other HBV components.

Accordingly, another aspect of the present invention contemplates a method for detecting an agent which exhibits inhibitory activity to an HBV which exhibits resistance or decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof,, said method comprising:

generating a genetic construct comprising a replication competent-effective amount of the genome from said HBV contained in a plasmid vector and then transfecting said cells with said construct;

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contacting said cells, before, during and/or after transfection, with the agent to be tested;

culturing said cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release virus or virus-like particles if resistant to said agent; and

subjecting the cells, cell lysates or culture supernatant fluid to viral-or viral-component-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of said agent.

Still another aspect of the present invention provides a method for detecting an agent which exhibits inhibitory activity to an HBV which exhibits resistance or decreased sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof, , said method comprising:

generating a genetic construct comprising a replication competent-effective amount of the genome from said HBV contained in or fused to an amount of a baculovirus genome effective to infect cells and then infecting said cells with said construct;

contacting said cells, before, during and/or after infection, with the agent to be 30 tested;

culturing said cells for a time and under conditions sufficient for the HBV to replicate, express genetic sequences and/or assemble and/or release, virus or virus-like particles if resistant to said agent; and

subjecting the cells, cell lysates or culture supernatant fluid to viral- or viral-component-detection means to determine whether or not the virus has replicated, expressed genetic material and/or assembled and/or been released in the presence of said agent.

Preferably, the HBV genome is stably integrated into the cells' genome.

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Particularly useful cells are 2.2.15 cells (Price et al., Proc. Natl. Acad. Sci. USA 86(21): 8541-8544, 1989 or AD cells (also known as HepAD32 cells or HepAD79 cells [Ying et al., Viral Hepat. 7(2): 161-165, 2000.

15 Whilst the baculovirus vector is a particularly useful in the practice of the present invention, the subject invention extends to a range of other vectors such as but not limited to adenoviral vectors.

The present invention further extends to cell lines (e.g. 2.2.15 or AD cells) carrying genetic constructs comprising all or a portion of an HBV genome or a gene or part of a gene therefrom.

The present invention also provides for the use of the subject HBV variants to screen for anti-viral agents. These anti-viral agents inhibit the virus. The term "inhibit" includes antagonizing or otherwise preventing infection, replication, assembly and/or release or any intermediate step. Preferred anti-viral agents include nucleoside or nucleotide analogs or anti-HBV agents, however, the present invention extends to non-nucleoside molecules.

In addition, rational drug design is also contemplated to identify or generate chemical molecules which either mimic a nucleoside or which interact with a particular nucleotide sequence or a particular nucleotide. Combinatorial chemistry and two hybrid screening are

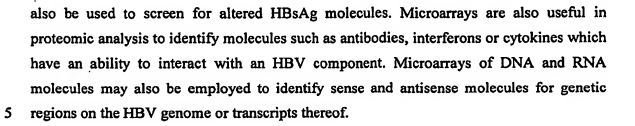
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some of a number of techniques which can be employed to identify potential therapeutic or diagnostic agents.

In one example, the crystal structure or the NMR structure of polymerase or the surface antigen is used to rationally design small chemical molecules likely to interact with key regions of the molecule required for function and/or antigenicity. Such agents may be useful as inhibitors of polymerase activity and/or may alter an epitope on the surface antigen.

- Several models of the HBV polymerase have been prepared due to the similarity with reverse transcriptase from HIV (Das et al., J. Virol. 75(10): 4771-4779, 2001; Bartholomeusz et al., Intervirology 40(5-6): 337-342 1997; Allen et al., Hepatology 27(6): 1670-1677, 1998). The models of the HBV polymerase can be used for the rational drug design of new agents effective against HBV encoding the resistant mutations as well as wild type virus. The rational drug that is designed may be based on a modification of an existing antiviral agent such as the agent used in the selection of the HBV encoding the mutations associated with resistance. Viruses or clones expressing HBV genomic material encoding the mutations may also be used to screen for new antiviral agents.
- In an alternative embodiment, the present invention also contemplates a method for detecting an agent which exhibits inhibitory activity to an HBV polymerase in an *in vitro* polymerase assay. The HBV polymerase activity can be examined using established assays (Gaillard et al., Antimicrob Agents Chemother. 46(4): 1005-1013, 2002; Xiong et al., Hepatology 28(6): 1669-1673, 1998).

As indicated above, microarray technology is also a useful means of identifying agents which are capable of interacting with defined HBV internal or external components. For example, arrays of HBV DNA polymerase or peptide fragments thereof carrying different amino acid-variants may be used to screen for agents which are capable of binding or otherwise interacting with these molecules. This is a convenient way of determining the differential binding patterns of agents between HBV variants. Arrays of antibodies may



The above methods are particularly useful in identifying an inhibitor of an HBV resistant to or exhibiting reduced sensitivity to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof. The present invention extends, therefore, to compositions of the inhibitors. The inhibitors may also be in the form of antibodies or genetic molecules such as ribozymes, antisense molecules and/or sense molecules for co-suppression or the induction of RNAi or may be other nucleoside or nucleotide analogs or other anti-HBV agents or derivatives of known analogs. Reference to RNAi includes reference to short, interfering RNAs (siRNA).

The term "composition" includes a "pharmaceutical composition" or a formulation.

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The inhibitor is referred to below as an "active ingredient" or "active compound" and may be selected from the list of inhibitors given above.

The composition may include an antigenic component of the HBV, a defective HBV variant or an agent identified through natural product screening or rational drug design (including combinatorial chemistry).

Pharmaceutically acceptable carriers and/or diluents include any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents and the like. The use of such media and agents for pharmaceutical active substances is well known in the art. Except insofar as any conventional media or agent is

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incompatible with the active ingredient, use thereof in the therapeutic compositions is contemplated. Supplementary active ingredients can also be incorporated into the compositions.

The pharmaceutical composition may also comprise genetic molecules such as a vector capable of transfecting target cells where the vector carries a nucleic acid molecule capable of encoding an aspartyl protease inhibitor. The vector may, for example, be a viral vector.

Pharmaceutical forms suitable for injectable use include sterile aqueous solutions (where water soluble) and sterile powders for the extemporaneous preparation of sterile injectable solutions. It must be stable under the conditions of manufacture and storage and must be preserved against the contaminating action of microorganisms such as bacteria and fungi. The carrier can be a solvent or dilution medium comprising, for example, water, ethanol, polyol (for example, glycerol, propylene glycol and liquid polyethylene glycol, and the like), suitable mixtures thereof and vegetable oils. The proper fluidity can be maintained, for example, by the use of superfactants. The preventions of the action of microorganisms can be brought about by various anti-bacterial and anti-fungal agents, for example, parabens, chlorobutanol, phenol, sorbic acid, thirmerosal and the like. In many cases, it will be preferable to include isotonic agents, for example, sugars or sodium chloride. Prolonged absorption of the injectable compositions can be brought about by the use in the compositions of agents delaying absorption, for example, aluminium monostearate and gelatin.

Sterile injectable solutions are prepared by incorporating the active compounds in the required amount in the appropriate solvent with the active ingredient and optionally other active ingredients as required, followed by filtered sterilization or other appropriate means of sterilization. In the case of sterile powders for the preparation of sterile injectable solutions, suitable methods of preparation include vacuum drying and the freeze-drying technique which yield a powder of active ingredient plus any additionally desired ingredient.

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When the active ingredient is suitably protected, it may be orally administered, for example, with an inert diluent or with an assimilable edible carrier, or it may be enclosed in hard or soft shell gelatin capsule, or it may be compressed into tablets. For oral therapeutic administration, the active ingredient may be incorporated with excipients and used in the form of ingestible tablets, buccal tablets, troches, capsules, elixirs, suspensions, syrups, wafers and the like. Such compositions and preparations should contain at least 1% by weight of active compound. The percentage of the compositions and preparations may, of course, be varied and may conveniently be between about 5 to about 80% of the weight of the unit. The amount of active compound in such therapeutically useful compositions is such that a suitable dosage will be obtained. Preferred compositions or preparations according to the present invention are prepared so that an oral dosage unit form contains between about 0.1 µg and 200 mg of active compound. Alternative dosage amounts include from about 1 µg to about 1000 mg and from about 10 µg to about 500 mg. These dosages may be per individual or per kg body weight. Administration may be per hour, day, week, month or year.

The tablets, troches, pills, capsules and the like may also contain the components as listed hereafter. A binder such as gum, acacia, corn starch or gelatin; excipients such as dicalcium phosphate; a disintegrating agent such as corn starch, potato starch, alginic acid and the like; a lubricant such as magnesium stearate; and a sweetening agent such as sucrose, lactose or saccharin may be added or a flavouring agent such as peppermint, oil of wintergreen or cherry flavouring. When the dosage unit form is a capsule, it may contain, in addition to materials of the above type, a liquid carrier. Various other materials may be present as coatings or to otherwise modify the physical form of the dosage unit. For instance, tablets, pills or capsules may be coated with shellac, sugar or both. A syrup or elixir may contain the active compound, sucrose as a sweetening agent, methyl and propylparabens as preservatives, a dye and a flavouring. Of course, any material used in preparing any dosage unit form should be pharmaceutically pure and substantially nontoxic in the amounts employed. In addition, the active compound(s) may be incorporated into sustained-release preparations and formulations.

As stated above, the present invention further extends to an isolated HBsAg from the HBV variants herein described. More particularly, the present invention provides an HBsAg or a recombinant form thereof or derivative or chemical equivalent thereof. The isolated surface component and, more particularly, isolated surface antigen or its recombinant, derivative or chemical equivalents are useful in the development of biological compositions such as vaccine formulations.

Yet another aspect of the present invention provides a composition comprising a variant HBV resistant to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or an HBV surface antigen from said variant HBV or a recombinant or derivative form thereof or its chemical equivalent and one or more pharmaceutically acceptable carriers and/or diluents. Such a composition may be regarded as a therapeutic composition and is useful in generating an immune response including a humoral response. Generally, the HBV variants are "defective" and in themselves are unable to cause a sustained infection in a subject.

- As indicated above, antibodies may be generated to the mutant HBV agents and used for passive or direct vaccination against infection by these viruses. The antibodies may be generated in humans or non-human animals. In the case of the latter, the non-human antibodies may need to be deimmunized or more specifically humanized prior to use. Deimmunized may include, for example, grafting complimentarity determining regions (CDRs) from the variable region of a murine or non-human animal anti-HBV antibody onto a human consensus fragment antibody binding (Fab) polypeptide. Alternatively, amino acids defining epitopes in the variable region of the antibody may be mutated so that the epitopes are no longer recognized by the human MHC II complex.
- 30 Insofar as ribozyme, antisense or co-suppression (RNAi) or siRNA or complexes thereof repression is concerned, this is conveniently aimed at post-transcription gene silencing.

DNA or RNA may be administered or a complex comprising RNAi or a chemical analog thereof specific for HBV mRNA may be employed.

All such molecules may be incorporated into pharmaceutical compositions.

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In another embodiment, the present invention provides a biological composition comprising a variant HBV or an HBsAg or L, M or S proteins from said variant HBV or a recombinant or derivative form thereof or its chemical equivalent.

10 Generally, if an HBV is used, it is first attenuated. The biological composition according to this aspect of the present invention generally further comprises one or more pharmaceutically acceptable carriers and/or diluents.

The biological composition may comprise HBsAg or like molecule from one HBV variant or the composition may be a cocktail of HbsAgs or L, M or S proteins or like molecules from a range of ADV- and/or LMV- and/or, FTC- and/or TFV-resistant HBV variants. Similar inclusions apply where the composition comprises an HBV.

The present invention is further directed to the use of defective HBV variants in the manufacture of therapeutic vaccines to vaccinate individuals against infection by HBV strains having a particular nucleotide sequence or encoding a particular polymerase or surface antigen or L, M or S proteins.

Examples of suitable vaccine candidates are defective forms of HBV variants comprising a mutation selected from, in one embodiment, in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another

embodiment includes rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, or a combination thereof or an equivalent mutation; in a further embodiment, sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, sI195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, or a combination thereof or an equivalent mutation.

In one embodiment, for example, an HBV variant may be identified having a particular mutation in its polymerase conferring resistance or decreased sensitivity to a nucleoside analog. This variant may then be mutated to render it defective, i.e. attenuated or unable to cause infection. Such a defective, nucleoside analog-resistant virus may then be used as a therapeutic vaccine against virulent viruses having the same mutation in its polymerase.

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The subject invention extends to kits for assays for variant HBV resistant to ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV. Such kits may, for example, contain the reagents from PCR or other nucleic acid hybridization technology or reagents for immunologically based detection techniques. A particularly useful assay includes the reagents and components required for immobilized oligonucleotide- or oligopeptide-mediated detection systems.

30 Still another aspect of the present invention contemplates a method for determining the potential for an HBV to exhibit reduced sensitivity to ADV, LMV, TFV, or FTC; or ADV

and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV and/or optionally other nucleoside or nucleotide analogs or other anti-HBV agents or combination thereof, said method comprising isolating DNA or corresponding mRNA from said HBV and screening for a mutation in the nucleotide sequence encoding HBV DNA polymerase resulting in at least one amino acid substitution, deletion and/or addition in any one or more of domains F and G, and domains A through to E or a region proximal thereto of said DNA polymerase and associated with resistance or decreased sensitivity to ADV, LMV, TFV, or FTC; or 10 ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV, wherein the presence of such a mutation is an indication of the likelihood of resistance to said ADV, LMV, TFV, or FTC; or ADV and LMV; ADV and TFV; LMV and TFV; FTC and ADV; FTC and TFV; 15 FTC and LMV; or ADV and LMV and TFV; or ADV and FTC and TFV; TFV and FTC and LMV; ADV and LMV and FTC; or ADV and FTC and LMV and TFV.

An assessment of a potential viral variant is important for selection of an appropriate therapeutic protocol. Such an assessment is suitably facilitated with the assistance of a computer programmed with software, which *inter alia* adds input codes for at least two features associated with the viral variants to provide a value corresponding to the resistance or sensitivity of a viral variant to a particular chemical compound or immunological agent. The I<sub>VS</sub> can be selected from (a) the ability to exhibit resistance for reduced sensitivity to a particular compound or immunological agent; (b) an altered DNA polymerase from wild-type HBV; (c) an altered surface antigen from wild-type HBV; or (d) morbidity or recovery potential of a patient. Thus, in accordance with the present invention, I<sub>VS</sub> for such features are stored in a machine-readable storage medium, which is capable of processing the data to provide a value for a particular viral variant or a biological specimen comprising same.

Thus, in another aspect, the invention contemplates a computer program product for assessing the likely usefulness of a viral variant or biological sample comprising same for determining an appropriate therapeutic protocol in a subject (Figure 3), said product comprising:

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(1) code that receives as input code for at least two features associated with said viral agents or biological sample comprising same, wherein said features are selected from:

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- (a) the ability to exhibit resistance for reduced sensitivity to a particular compound or immunological agent;
- (b) an altered DNA polymerase from wild-type HBV;
- (c) an altered surface antigen from wild-type HBV; or
- 15 (d) morbidity or recovery potential of a patient;
  - (2) code that adds said input code to provide a sum corresponding to a value for said viral variants or biological samples; and
- 20 (3) a computer readable medium that stores the codes.

In a related aspect, the invention extends to a computer for assessing the likely usefulness of a viral variant or biological sample comprising same in a subject, wherein said computer comprises:

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(1) a machine-readable data storage medium comprising a data storage material encoded with machine-readable data, wherein said machine-readable data comprise input codes for at least two features associated with said viral variant or biological sample; wherein said features are selected from:-

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- (a) the ability to exhibit resistance for reduced sensitivity to a particular compound or immunological agent;
- (b) an altered DNA polymerase from wild-type HBV;
- (c) an altered surface antigen from wild-type HBV; or
- (d) morbidity or recovery potential of a patient;
- (2) a working memory for storing instructions for processing said machinereadable data;
- 10 (3) a central-processing unit coupled to said working memory and to said machine-readable data storage medium, for processing said machine readable data to provide a sum of said input code corresponding to a value for said compound(s); and
- 15 (4) an output hardware coupled to said central processing unit, for receiving said value.

Any general or special purpose computer system is contemplated by the present invention and includes a processor in electrical communication with both a memory and at least one input/output device, such as a terminal. Figure 3 shows a generally suitable computer system. Such a system may include, but is not limited, to personal computers, workstations or mainframes. The processor may be a general purpose processor or microprocessor or a specialized processor executing programs located in RAM memory. The programs may be placed in RAM from a storage device, such as a disk or pre-programmed ROM memory. The RAM memory in one embodiment is used both for data storage and program execution. The computer system also embraces systems where the processor and memory reside in different physical entities but which are in electrical communication by means of a network.

30 In an alternative embodiment, the program screens for a mutation selected from, in one embodiment, in one embodiment, rtT38K, rtR55H and rtA181V; in another embodiment

includes rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtE142V, rtA/T181A rtI204M, rtQ/P/S/Stop215S, rtE/K218E, rtN/H238H and rtY245H; in yet another embodiment includes rtN238T; and yet another embodiment includes rtI122V and rtA181T; in yet another embodiment includes rtL180M, rtA/V200V, rtM204V, rtV214A, rtH237H/P, rtV253G, in yet another embodiment includes rtT128N and rtN236T, in yet another embodiment includes tL180M, rtM204V rtQ215S, in yet another embodiment includes rtT128S rtL180M, rtM204Vand rtQ215S, in yet another embodiment includes rtI80L, rtI204M, rtN238T, in yet another embodiment includes rtN238T/A, in yet another embodiment includes rtI187V, or a combination thereof or an equivalent mutation; in a further embodiment, sQ30K. sE44G, sA47T, sI126T, sA159V and sL173F, in another embodiment include sF55S, sC/Stop69C, sC/Y76Y, sI/V110I, sN/T131N, sN134Y, sStop/W172W, sStop/W196W, sS/R207R; in yet another embodiment include sV14A, sL95W, sV96G, and sI208T/I; and in yet another embodiment include sT47A and sW172stop, and in yet another embodiment include PreS2 T6S, sT47A, sP62L, sL/F192F, s1195M, and in yet another embodiment include sS53L, and in yet another embodiment include sP120T, and in yet another embodiment include sN40S, sS207R, and in yet another embodiment include sQ101R, sI195M, sS207R, and in yet another embodiment include sL95W and sL196W, and in yet another embodiment include sV14A, or a combination thereof or an equivalent mutation.

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The present invention is further described by the following non-limiting Examples.

### **EXAMPLE 1**

### Overlapping genome of HBV

The overlapping genome of HBV is represented in Figure 1. The gene encoding DNA polymerase (P), overlaps the viral envelope genes, Pre-S1 and Pre-S2, and partially overlaps the X and core (C) genes. The HBV envelope comprises small, middle and large proteins HBV surface antigens. The large protein component is referred to as the HBV surface antigen (HBsAg) and is encoded by the S gene sequence. The Pre-S1 and Pre-S2 gene sequences encode the other envelope components.

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#### **EXAMPLE 2**

# Patients on ADV Treatment and Analysis of HBV DNA

Patient A: During ADV treatment, unique HBV mutations were detected by sequencing (Table 4). This includes the unique mutation at rtT38K, and rtA181V. A number of other changes were also detected in the polymerase rtR55H and in the overlapping envelope gene (Table 4, Figures 4, 5 and 6). The changes in the HBsAg include sQ30K, sE44G, sA47T, sI126T, sA159V and sL173F. These unique changes were compared to reference sequences from each of the seven genotypes A-G as well as a consensus sequence from pretreatment samples to determine unique changes

The HBV mutations during ADV treatment are listed in Table 5 and Figures Patient B: 7, 8, and 9. The unique changes in the rt region of the HBV DNA polymerase include Other changes in the HBV polymerase while on ADV treatment include rtY245H. rtE142V, rtA/T181A rtI204M, rtS/T78S, rtV80L, rtN/S118N, rtN/K139K, rtQ/P/S/Stop215S,rt E/K218E, rtN/H238H. The changes in the HBsAg while on ADV sI/V110I, sN/T131N, sN134Y, treatment include sF55S, sC/Stop69C, sC/Y76Y, sStop/W172W, sStop/W196W, sS/R207R.

30 Patient C: The HBV mutations prior to ADV treatment and during ADV treatment are listed in Table 6 and Figures 10, 11 and 12. The unique changes in the rt region of the

HBV DNA polymerase while on ADV treatment include rtN238T. The unique changes in the HBsAg include sV14A, sL95W, sV96G, and sI208T/I

Patient D: The HBV mutations during ADV treatment is listed in Table 7 and Figures 13, 14 and 15. The unique changes in the HBV DNA polymerase include rtI122V and rtA181T. The unique changes in the surface include sT47A and sW172stop.

Patient E. This patient was previously treated with lamivudine and selected the unique mutations rtH237H/P while on LMV. This patient did not respond to ADV treatment 10 Changes in the polymerase on ADV treatment include rtL180M, rtA/V200V, rtW204V, rtV214A, rtH237H/P and rtV253G. The unique changes in the surface include PreS2 T6S, sT47A, sP62L, sL/F192F, sI195M. NB: Patient on TFV and responded Changes on ADV listed in Table 8 and Figures 16, 17 and 18.

Patient F: Unique changes during ADV treatment include the polymerase mutations at rtN238T and envelope mutations at sS53L. Changes on ADV listed in Table 9 and Figures 19, 20 and 21.

Patient G: Unique mutations while on ADV treatment include changes in the polymerase rtT128N and rtN236T and a change in envelope sP120T. Changes on ADV listed in Table 10 and Figures 22, 23 and 24.

Patient H: Mutations in the polymerase gene while on ADV treatment include rtL180M, rtM204V rtQ215S. Changes in envelope gene includesN40S, sS207R. Changes on ADV listed in Table 11 and Figures 25, 26 and 27.

Patient I: Mutations in the polymerase gene while on ADV treatment include rtT128S rtL180M, rtM204Vand rtQ215S, while mutations in the envelope gene included sQ101R, sI195M, sS207R. Changes on ADV listed in Table 12 and Figures 28, 29 and 30.

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- Patient J: Mutations in the polymerase gene included rtI80L, rtI204M, rtN238T and mutations in envelope sL95W and sL196W during ADV treatment. Changes on ADV listed in Table 13 and Figures 31, 32 and 33.
- 5 Patient K: Mutations in the polymerase gene at rtN238T/A was detected during Adv treatment. No changes in envelope were detected during treatment. Changes on ADV listed in Table 14 and Figures 34, 35 and 36.
- Patient L: Mutations in the polymerase gene at rtI187V was detected during ADV treatment. A mutation in the envelope gene at sV14A was also detected. Changes on ADV listed in Table 15 and Figures 37, 38 and 39.

# **EXAMPLE 3**

# **Detection of Viral Markers**

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Hepatitis B surface antigen (HBsAg), hepatitis B e antigen (HBeAg), anti-HBe and hepatitis B core antigen (HBcAg) specific IgG and IgM were measured using commercially available immunoassays (Abbott Laboratories, North Chicago, IL, USA). Hepatitis B viral DNA levels were measured using a capture hybridization assay according to the manufacturer's directions (Digene Hybrid Capture II, Digene Diagnostics Inc., Beltsville, MD). The manufacturers stated cut-off for detecting HBV viremia in clinical specimens was 0.7x10<sup>6</sup> copies/ml or 2.5 pg/ml, [Hendricks et al., Am J Clin Pathol 104: 537-46, 1995]. HBV DNA levels can also be quantitated using other commercial kits such as Cobas amplification HBV monitor kit (Roche).

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#### **EXAMPLE 4**

# Sequencing of HBV DNA

HBV DNA was extracted from 100 µl of serum as described previously by Aye et al., J. 30 Hepatol. 26: 1148-1153, 1997. Oligonucleotides were synthesized by Geneworks,

Adelaide, Australia. Amplification of the HBV polymerase gene has been described by Aye et al., 1997, supra.

The specific amplified products were purified using PCR purification columns from MO

BIO Laboratories Inc (La Jolla, CA) and directly sequenced using Big Dye terminator
Cycle sequencing Ready Reaction Kit (Perkin Elmer, Cetus Norwalk, CT). The PCR
primers were used as sequencing primers, OS1 5'- GCC TCA TTT TGT GGG TCA CCA
TA-3' (nt 1408-1430) [SEQ ID NO:3], TTA3 5'-AAA TTC GCA GTC CCC AAA3'(nt2128-2145) [SEQ ID NO:4], JM 5'-TTG GGG TGG AGC CCT CAG GCT 3'(nt1676-1696) [SEQ ID NO:5], TTA4 5'-GAA AAT TGG TAA CAG CGG -3' (nt 26152632) [SEQ ID NO:6], OS2 5' TCT CTG ACA TAC TTT CCA AT 3' (nt 2798-2817)
[SEQ ID NO:7], to sequence the internal regions of the PCR products.

#### **EXAMPLE 5**

#### Adefovir Dipivoxil (ADV)

ADV (formerly Bis-pom PMEA)) is a potent inhibitor of HBV replication. The structure of ADV is shown in Figure 2 and its synthesis is described by Benzaria *et al.*, *J Med Chem.* 39: 4958-4965, 1996).

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Those skilled in the art will appreciate that the invention described herein is susceptible to variations and modifications other than those specifically described. It is to be understood that the invention includes all such variations and modifications. The invention also includes all of the steps, features, compositions and compounds referred to or indicated in this specification, individually or collectively, and any and all combinations of any two or more of said steps or features

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Table 4: Patient A HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV viral load	HBV	RT	HBsAg Mutations
			Mutations		
ADV	22/10/02	8.77E+06	•		sT47A sT131T/I
ADV	14/01/03	1.21 E+09	•		-
ADV	8/7/03	7.92 E+07	rtT38K rtA181V		sL173F

Table 5: Patient B RT and Polymerase mutations detected during ADV therapy

Treatment	Date	HBV RT Mutations	HBsAg Mutations
ÁDV	13/02/03	rtV80L - rtN118N/S rtN139N/K - rtV142E rtA181A/T rtI204M rtQ/P/S/Stop215Q rtE218K/E rtN238N/H	sC76Y/C sI110V/I sN131N/T sY134N sW172Stop/W sStop196W sR/S207S
ADV	21/03/03	rtS78T/S rtV80L - rtN118N/S rtN139N/K - rtV142E rtA181A/T rtI204M rtQ215 Q/P/S/Stop rtE218K/E rtN238N/H	sS55F sC69Stop - sC76Y/C sI110V/I sN131N/T - sY134N sW172Stop/W sStop/W196W sS207S/R

ADV	22/07/03	rtS/T78S	sF55S
		rtV80L	`sC/Stop69C
		rtN/S118N	sC/Y76Y
		rtN/K139K	sI/V110I
1		-	sN/T131N
		rtE142V	-
		rtA/T181A	sN134Y
Į.		-	sStop/W172W
]	<b>\</b>	rtI204M	sStop/W196W
		rtQ/P/S/Stop21	sS/R207R
		rt5SE/K218E	
		rtN/H238H	
		rtY245H	

Table 6: Patient C HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations
		Mutations		
ADV	18/08/03	rtN238T		sV14A
				sL95W
				sV96G
				sI208T/I

Table 7: Patient D HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations
		Mutations		
ADV	20/08/03	rtI122V		sT47A
		rtA181T		sW172Stop

Table 8: Patient E HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV RT Mutations	HBsAg Mutations
LMV	23/05/02	rtL180M rtA200V/A rtM204V rtV214A rtP237H	sL192L/F sI195M
ADV	17/07/03	rtL180M rtA/V200V rtM204V rtV214A rtH237H/P rtV253G	PreS2 T6S sT47A sP62L sL/F192F s1195M

Table 9: Patient F HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations
		Mutations		
ADV	16/10/03	rtN238T		sS53L

Table 10: Patient G HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV Mutations	RT	HBsAg Mutations
ADV	12/11/03	rtT128N rtN236T		sP120T

Table 11: Patient H HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations
		Mutations		
ADV	05/11/03	rtL180M rtM204V rtQ215S		sN40S sI195M sS207R

Table 12: Patient I HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations
		Mutations		
ADV	05/02/03	rtT128S rtL180M rtM204V rtQ215S	,	sQ101R - - sI195M - sS207R

Table 13: Patient J HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations	
		Mutations			
ADV	18/11/03	rtI80L - rtI204M rtN238T		sL95W sL196W -	

Table 14: Patient K HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations
		Mutations		
ADV	31/03/03	rtN238T/A		No changes

Table 15: Patient L HBV Polymerase and envelope mutations detected during ADV therapy

Treatment	Date	HBV	RT	HBsAg Mutations
		Mutations		
ADV	17/09/03	rtI187V		sV14A



1 of 51

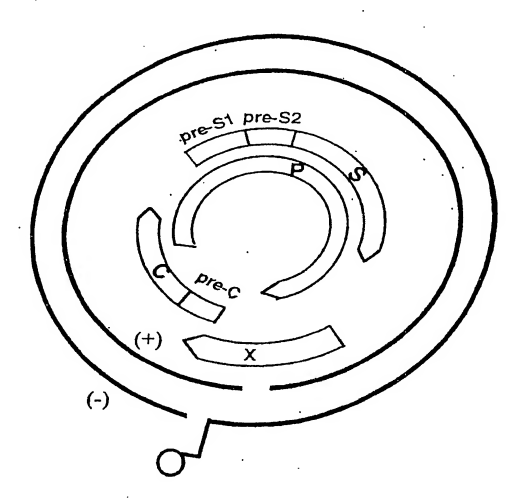


Figure 1

Figure 2



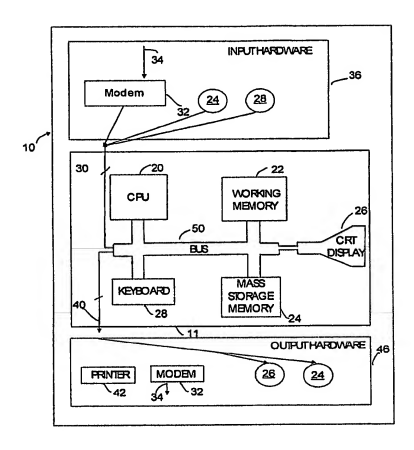


Figure 3A



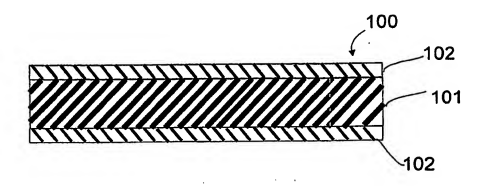


Figure 3B

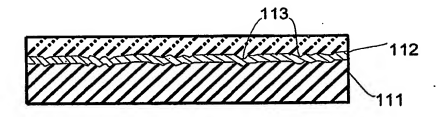


Figure 3C

Figure	4:	Patient	A nt	sequence	•		
-			10	20		40	50
		GCTTCC	CACCAP	ATCGGCAGGC	AGGAAGACAGC	CTACTCCCAT	CTCTCCACC
			60	70	. 80	90	100
		TCTAAG			AGGCCATGCAG		
				100	100	140	150
		አርርአምር	110	120	130 AGACCTGCTGG	140 TGGCTCCAG	150
		ACCATO	SCICIO	SCIAGAICCC	AGACCIGCIGG	100C1CCAG	CCGGHACA
•			160	170		190	200
		GTAAAC	CCTGI	TCCGACTAC	rgcctctccca	TATCGTCAAT	CTTCTCGAG
			210	220	230	240	250
		GACTG			ATATGGAGAGC		
			260	270			300
		GACCC	CTGCTC	CGTGTTACAG	GCGGGGTTTTI	CTTGTTGAC	AAGAATCCTC
			310	320	330	340	350
		ACAATA			CTCGTGGTGGA		ATTTTCTAGG
			360	370		390	
		GGGAGG	CACCCA	ACGTGTCCTG	GCCAAAATTTG	CAGTCCCCA	ACCICCAAIC
			410	420	430	440	450
		ACTCAC			CCAATTTGTCC	TGGTTATCG	CTGGATGTGT
		000000	460		480	490	
		CTGCGC	oCGTT1	PTATCATCTT	CCTCTTCATCC	.IGCIGCIAI	SCCICATOIT
			510	520	530	540	550
		CTTGTT	CGGTT	CTTCTGGACT.	ACCAAGGTATO	TTGCCCGTT'	TGTCCTCTAC
			E 6 0	570	580	590	600
		<b>ጥጥርር Δ</b> (	560 מסממני	_	AGCACGGGACO		
		1100/10	3011101	11 0111011100			
			610	620		640	
		CCTGC	rcaag(	GAACCTCTAT	GTTTCCCTCTT	GTTGCTGTA	CAAAACCTTC
			660	670	680	690	700
		GGACG			TTCCCATCCC		
		an mme	710	720		740	750
		GATTC	_TATG(	3GAGTGGGCC	TCAGTCCGTTT	CTCCTGGTT	CAGTTTACTA
			760	770	780	790	800
		GTGCC			CGTAGGGCTTT		

	810	820	830	840	850
AGTTA	ratggatg	ATGTGGTATT	GGGGCCAAG	rctgtacaac.	ATCTTGA
		•			
	860	870	880	890	900
ATCCC	COATATT	GCTATTACCA	TTTTCTTTTC	STCTTTGGGT.	ATACATT
	910	920	0 <i>É</i> e	940	950
TAAAC		AACCAAGCGTT		CCCTTAACTT	CATGGGA
	960	970	980	990	1000
TATGT		GTTGGGGTAC			
AATCA	A A				

Figure 4

#### Figure 5: Patient A. HBV Polymerase sequence

10	20	30	40	50
EDWGPCAEYGEHHI	RIPRTPARVI	GGVFLVDKNI	PHNTKESRLV	/DFSQFS
60	70	80	90	100
RGSTHVSWPKFAVP	NLQSLTNLLS	SULSMISID	/SAAFYHLPLI	IPAAMPH
110	120	130	140	150
LLVGSSGLPRYVAR	LSSTSRNINY	QHGTMQDLH	DSCSRNLYVSI	LLLYKT
160	170	180	190	200
FGRKLHLYSHPIIL			rsaicsvvrr	AFPHCLA
210	220	230	240	250
FSYMDDVVLGAKSV				
260	270			
GYVIGSWGTLPQEH				

#### Figure 6: Patient A HBV HbsAg sequence

10	20	30	40	50		
MESTTSGFLGPLLV	LQAGFFLLTE	RILTIPKSLDS	SWWTSLNFLGG	SAPTCPG		
60	70	80	. 90	100		
QNLQSPTSNHSPTS	SCPPICPGYRV	WCLRRFIIF	LFILLLCLIFI	TATTDA		
110	120	130	140	150		
QGMLPVCPLLPGTS	STTSTGPCKT	CTTPAQGTSMI	FPSCCCTKPSI	GNCTCI		
160	170	180	190	200		
PIPSSWAFVRFLWEWASVRFSWFSLLVPFVQWFVGLSPTVWLSVIWMMWY						
210	220					
WGPSLYNILNPFI	PLLPIFFCLWY	/YI				

#### Figure 7: Patient B HBV NT sequence

10	20		40	50
TCTGTCTCCAC	CCTTTGAGAGA		CAGGCCATGC	AGTGGAACT
· 60	70	80	90	100
CCACAACCTTO	CCACCAAACTC	TGCAAGATCC	CCAGAGTGAGA	GGCCTGTAT
110		130	140	150
TTCCCTGCTG		TCAGGAACAG	STAAACCCTGT	TCCGACTTC
160	170	180	190	200
TGTCTCTCAC	ACATCGTCAAT	CTTCTCGAG	GATTGGGGTCC	CTGCGCTGA
210 ACATGGAGAAC		230 GATTCCTAGE		250 GTGTTACAG
260	270	280	290	300
GCGGGGTTTT	CTTGTTGACA	AGAATCCTC	ACAATACCGCA	GAGTCTAGA
310 CTCGTGGTGG		330 ATTTTCTAGG		350 TGTGTCTTG
360	370	380	390	400
GCCAAAATTC	GCAGTCCCCA	ACCTCCAATC	ACTCACCAACC	TCCTGTCCT
410 CCAACTTGTC			440 CTGCGGCGTTT	450 TATCATCTT
460	470	480	490	500
CCTCTTCATC	CTGCTGCTATO	SCCTCATCTT	CTTGTTGGTTC	TTCTGGACT
510 ATCAAGGTATO		530 GTCCTCTAA		550 TCAACCACC
560	570	580	590	600
AGCACGGGAC	CATGCAGAACO	CTGCACGACT	CCTGCTCAAGG	AAACTCTAT
610	620	630	640	650
GTATCCCTCC	IGTTGCTGTAC	CCAAACCTTC	GGACGGAAATT	GCACCTGTA
660	670	680	690	700
TTCCCATCCC	ATCATCCTGG	CTTTCGGAA	AATTCCTATGG	GAGTGGGCC
710	720	730	740	750
TCAGCCCGTT	CTCCTGGCT	CAGTTTACTAC	GTGCCATTTGT	TCAGTGGTT

•	760	770	780	790	800
CGTAGGG	CTTTCCCCC	ACTGTTTGGCT	TTTCAGTTATA	ATGGATGATGT	GGT
8	310	820	830	840	850
ATTGGGG	SCCAAGTCT	STATCGCATC1	TTGAGTCCCT	TTTACCGCTG	TTA
1	860	870	880	890	900
CCAATTT	PCTTTTGTCT	TTTGGGTATAC	CATTTAAACC	CTCACAAAACA	AAA
	910	920	930	940	950
AGATGGG	GTCACTCTT	PACATTTCATO	GGCTATGTC	ATTGGATGTTA	TGG
•	960	970	980		
GTCATTG	CCACAAGAT	CACATCAGACA	AGAAAA		

Figure 7 continued

#### Figure 8: Patient B POLYMERASE sequence

	10	20	30	40	50
EDWGPCAEI	HGEHHIRIP	RTPARVTGGV	/FLVDKNPHN1	PAESRLVVDFS	SQFS
	60	70	80	90	100
RGNYRVSWI	PKFAVPNLQ	SLTNLLSSNI	LSWLSLDVSAA	AFYHLPLHPA <i>I</i>	HAW
1:	10	120	130	140	150
LLVGSSGLS	SRYVARLSS	NSRIFNHQHO	STMQNLHDSCS	RKLYVSLLLI	LYQT
16	60	170	180	190	200
FGRKLHLY:	SHPIILGFR	KIPMGVGLSE	FLLAQFTSAI	CSVVRRAFP	ICLA
2:	10	220	230	240	250
FSYMDDVVI	LGAKSVSHL	ESLFTAVTNE	FLLSLGIHLNE	PHKTKRWGHS	LHFM
20	60				
GYVIGCYG	SLPQDHIRQ	K			

#### Figure 9:Patient B HBsAG sequence

10	20	30	40	50
MENITSGFLGP	LLVLQAGFFL	LTRILTIPQS	LDSWWTSLNF	LGGTTVCLG
	_			
· 60	70	. 80	90	100
QNSQSPTSNHS	PTSCPPTCPG	YRWMYLRRFI	IFLFILLLCL	IFLLVLLDY
				150
110	120	130	140	150
QGMLPVCPLIP	GSSTTSTGPC	RTCTTPAQGN	ISMYPSCCCTK	PSDGNCTCI
160	. 170	180	190	200
PIPSSWAFGKF				
210	220			
WGPSLYRILSP	FLPLLPIFFC	LWVYI		

#### Figure 10: Patient C HBV NT sequence

CAGCAGO	10 GCCTCCTCC	20 IGCCTCCTCC	30 AATCGGCAGT	40 CAGGAAGACA	50 GCCT
ACTCCCA	60 ATCTCTCCAC	70 CTCTAAGAGA	80 CAGTCATCCT	90 CAGGCCATGC	
1.01000					
GAACTCC	110 CAGCACATTC	120 CACCAAGCTC	130 TGCTAGATCC	140 CAGAGTGAGG	150 GGCC
	160.	170	180	190	200
TATATTI			TCCGGAACAG		
	210	220	230	240	250
ACTACTO			CTTCTCGAGG.		
N C C C N N C	260	270 ACCACAMCAC	280 GATTCCTAGG	290 <b>»</b> ССССТЕСТС	300 CCCT
ACCGAAC	AIGGAGAGC	ACCACAT CAG	GATICCIAGO	ACCCCIGCIC	0001
	310	320	330	340	
TACAGGO	CGGGGTTTTT	CTTGTTGACA	AGAATCCTCA	CAATACCACA	GAGT
	360	370	380	390	400
CTAGACT			TTTTCTAGGG		
	410	420	430	440	450
TCCTGGC			CCTCCAATCA		
	460	470	480	490	500
GTCCTCC	CAATTTGTCC	TGGTTATCGC	TGGATGTGTC	TGCGGCGTTT	TAIC
	510 .		530		
ATCTTC	CTCTTCATCC	TGCTGCTATG	CCTCATCTTC	TTGTGGGGTC	TTCT
	560	570	580	590	600
GGACTA			GTCCTCTACT		
ርሞልርሮል(			630 TGCACGACTC		
OINOON					
	660	670	68 <u>.</u> 0	690	700
TCTATG	TTTCCCTCTT	GTTGCTGTAC	AAAACCTTCG	GACGGAAATI	GCAC

Figure 10

710	720	730	740	750
TTGTATTCCCATCC	CATCATCTT	GGGCTTTCGC	AGATTCCTAT	rgggagt
			790	800
GGGCCTCAGTCCGT	TTCTCCTGG	CTCAGTTTACT	AGTGCCATT	rgttcag
810			840	850
TGGTTCGTAGGGCT	TTCCCCCAC'	<b>IGTTTGGCTT</b> 1	TAGTTATAT	GGATGAT
860 .	870	880	890	900
GTGGTATTGGGGGC			SAATCCCTTT	TACCGC
910	920	930	940	950
TGTTACCAATTTTC	TTTTGTCTT	TGGGTATACAT	TTAAACCCT	ACTAAAA
960	970	980	990	1000
CCAAACGTTGGGGC				
1010	1020	1030	1040	
TGGGGTACCTTACC	ACAAGAACA	TATTGTACAC	AAAATCAGAC	AA

Figure 10 continued

#### Figure 11: Patient C Polymerase sequence

1	0	20	30	40	50
EDWGPCTEH	GEHHIRIP:	RTPARVTGGV	FLVDKNPHNI	TESRLVVDFS	SQFS
6	0	70	80	90	100
RGNTQVSWP	KFAVPNLQ	SLTNLLSSNI	SWLSLDVSAA	YEATHT THE STATE OF THE STATE O	MPH
11	0	120	130	140	150
LLVGSSGLP	RYVARLSS	TSRNINYQHO	STMQDLHDSCS	SRNLYVSLLLI	LYKT
16	0	170	180	190	200
FGRKLHLYS	HPIILGFR	KIPMGVGLSE	PFLLAQFTSA	CSVVRRAFPI	ICLA
21	-		230	240	250
FSYMDDVVL	GAKSVQHL	ESLFTAVTNE	FLLSLGIHLNI	PTKTKRWGYS	LNFM
				•	
26	0	270			
GYVIGSWGT	LPQEHIVH	KIRQ			

Figure 11

#### Figure 12 Patient C HbsAg sequence

	10	20	30	40	50
MESTTS	FLGPLLALO	AGFFLLTRIL'	TIPQSLDSWW	TSLNFLGGTP	KCPG
			<b>~</b>		
	60	70	0.0	90	100
	60	70	80	• •	
ONLOSPI	SNHSPTSCP:	PICPGYRWMC	LRRFIIFLFI	LLLCLIFLWG	LLDY
<b>2</b> , - · · · · · · · · · · · · · · · · · ·					
	110	120	130	140	150
	110				
QGMLPV	CPLLPGTSTT	STGPCKTCTT	PAQGTSMFPS	CCCTKPSDGN	CTCI
•	160	170	180	190	200
PIPSSW	AFARFLWEWA	SAKERMTZTT	VPFVQWFVGL	SLIAMTTAIM	MMM I
				·	
	210	220			
MODGE W		DY BBOT MINT			
MGBSTI	NXLNPFLPLL	RICECTMAIT			

Figure 12

#### Figure 13; Patient D NT sequence

10 CTCCTGCATC	20 PACCAATCGG		40 GACAGCCTACT	
60 CCACCTCTAA			90 ATGCAGTGGAA	100 ACTCCACAAC
110	120		140	150
TTTCCACCAA	GCTCTGCTAG		GAGGGGCCTCT	ATTTTCCTG
160 CTGGTGGCTC	170 CAGTTCCĠGG			200 ACTGCCTCT
210	220	230		250
CCCATATCGT	CAATCTTCTCC	SAGGACTGGG		GAACATGGA
260	270		290	300
GAGCACAACA	CAGGATTCC		GCTCGTGTTAC	AGGCGGTGT
310	320		340	350
TTTTCTTGTT	GACAAGAATCO		CACAGAGTCTA	AGACTCGTGG
360	370		390	400
TGGACTTCTC	CAATTTTCT		CCCGCGTGTC	CTGGCCAAAA
410	420	430		450
TTCGCAGTCC	CCAACCTCCA	ATCACTCACC		CCTCCAATTT
460	470	480		500
GTCCTGGCTA	CGCTGGATG	FGTCTGCGGC		CTTCCTCTTC
510 ATCCTGCTGC			540 GTTCTTCTGG	550 ATTACCAAGG
560 TATGTTGCCC			590 AACGTCAACT	
610 GACCATGCAA		630 ATTCCTGCTC		650 PATGTTTCCC
660	670	680	690	700
TCATGTTGCT	GTACAAAACC	TTCGGACGGA	AACTGCACTT	STATTCCCAT
710	720	730	740	750
CCCATCATCC	TGGGCTTTCG	CAAGATTCCT	ATGGGAGTGG	SCCTCAGTCC

Figure 13

760	770	780	790	800
GTTTCTCTTGACTCAG	TTTACTAGT	GCCATTTG	TTCAGTGGTTC	TAGGG
810	820	830	840	850
CTTTCCCCCACTGTTT	GGCTTTCAG	TTATATGG	ATGATGTGGTAT	TGGGG
860	870	880	890	900
GCCAAGTCTGTACAAC	ATCTTGAGT	CCCTTTAT	ACCGCTATTAC	CAATTT
910	920	930	940	950
TCTTTTGTCTTTGGGT	ATACATTTA	AACCCTAA	TAAAACCAAGC	SATGGG
960	970	980	990	1000
GTTACTCCCTTAACTT	CATGGGATA	TGTCATTG	GAAGTTGGGGG	ACTTTA
1010	1020			
CCACAGGAACATATTO	STGCTC			

Figure 13 continued

#### Figure 14: patient D HBV POL sequence

	10	20	30	40	50
EDWGPCTE	EHGEHNIRI	PRTPARVTG	SVFLVÖKNPHN	TTESRLVVI	DFSQFS
	60	70	80	90	100
RGSTRVS	VPKFAVPNL(	QSLTNLLSSN	ILSWLSLDVSA	AFYHLPLHI	PAAMPH
	L10	~~~	130	140	150
LLVGSSG	LPRYVARLS	STSRNVNYQ	IGTMQDLHDSC	SRNLYVSL	<b>1LLYKT</b>
	160	170	180	190	200
FGRKLHL	YSHPIILGF	RKIPMGVGLS	SPFLLTQFTSA	ICSVVRRAI	FPHCLA
	210		230	240	250
FSYMDDV	VLGAKSVQHI	LESLYTAIT	NFLLSLGIHLN	PNKTKRWG	YSLNFM
-	260	•			
GYVIGSW	GTLPQEHIV	L			

Figure 14



#### Figure 15 Patient D HBsAg sequence

	10	20	30	40	50
MESTTSGE	LGPLLVLQA	VFFLLTRILT	IPQSLDSWW:	rslnflgeap?	<b>ACPG</b>
	_				
	60	70	80	90	100
QNSQSPTS	SNHSPTSCPI	PICPGYRWMCL	RRFIIFLFI	LLLCLIFLLV	LLDY
•	110		130	140	150
QGMLPVC	PLLPGTSTTS	STGPCKTCTIP	AQGTSMFPS(	CCCTKPSDGN	CTCI
-	160		180	190	200
PIPSSWA	FARFLWEWAS	SVRFS*LSLLV	PFVQWFVGL	SPTVWLSVIW	YWMN
2	210	220			
WGPSLYN:	ILSPFIPLLI	PIFFCLWVYI			

Figure 16:	Patient	E HBV nt	sequence		
10		0	30	40 TTCCACCAAGO	50 TCT
AGTCATCCTC	AGGCCATGC	AGTGGAAC	TCCAGCACAI	LICCACCAAGC	,101
60 GCTAGATCC		0 GGCCTATA		90 rggtggctcc <i>p</i>	100 GTT
110			130	140	150
CAGGAACAGT	AAACCCTGT	TCCGACTA	CTGCCTCTCC	CCATATCGTCA	MIC
160			180		200
TTCTCGAGG	CTGGGGACC	CTGCACCG	AATATGGAG <i>I</i>	AGCACCACATO	AGG
210			230		250
ATTCCTAGGA	ACCCCTGCTC	GTGTTACA	GGCGGGGTT	TTTCTTGTTG <i>I</i>	ACAA
		•	000	200	200
260			280		300 ייממי
GAATCCTCAC	CAATACCACA	GAGTCTAG	ACICGIGGI	GGACTTCTCT(	NUL
310	32	:0	330	340	350
				TTGCAGTCCC	CAAC
360			380		400
CTCCAATCAC	CTCACTAACC	TCTTGTCC	TCCAATTTG!	rcctggttat(	CGCT
444	. 40		420	440	450
410				rcctgctgct/	
GGAIGIGIC.	160666111	INICAICI	. ICCICITON	1001001001.	
460	) 47	0	480	490	500
CTCATCTTC			CTACCAAGGT	ATGTTGCCCG'	TTTG
510			530		550
TCCTCTACT'	rccaggaac <i>i</i>	ATCAACTAC	CAGCACGGG	ACCATGCAAG	ACCT
56	0 57	7.0	580	590	600
				CTTGTTGTTG'	
00710071010	010010.11.00				
61			630	640	650
AAACCTTCG	GACGGAAATI	GCACTTG	TATTCCCATC	CCATCATCTT	GGGC
66	0 67	70	680	690	700
				TTTCTCATGG	CTCA

Figure 16

710	720	730	740	750	
GTTTACTAGT	CCATTTGTTCA	GTGGTTCGT	AGGGCTTTCC	CCCACTGTT	
				•	
760	770	780	790	800	
TGGTTTTCAGT	TATGTGGATGA	TGTGGTATT	GGGGCCAAG	<b>PCTGCACAA</b>	
810	820	830	840	850	
CATCTTGAATC	CCTTTTTACCG	CTATTACCA	ATTTTCTTTT(	GTCTTTGGG	
860	870	880	890	900	
TATACATTTA	ACCMTAATAAA	ACCAAACGT	rggggctatt	CCCTTAACT	
910	920	930	940	950	
TTATGGGATATGGAATTGGAAGTTGGGGTCCTGCCCAGGGAAGATGGCAG					
GGG					

Figure 16 continued

Figure 17 Patien	t E: HBV	polymerase		
10		30		50
SSSGHAVELQHIPPS	SAKSQSEGE	TPSCMMTGEK	NSKPCSDICI	POUTAND
60		80	90	100
LEDWGPCTEYGEHHI	RIPRTPARV	TGGVFLVDKN	PHNTTESRLV	/VDFSQF
		130		150
SRGSTRVSWPKFAVP	NLQSLTNLI	SSNLSWLSLD	VSAAFYHLPI	LHPAAMP
		180		200
HLLVGSSGLPRYVAR	LSSTSRNIN	IYQHGTMQDLH	DSCSRNLYVS	SLLLLYK
		230		250
TFGRKLHLYSHPIIL	GFRKI PMG\	/GLSPFLMAQF	TSAICSVVRI	RAFPHCL
			290	300
VFSYVDDVVLGAKSA	QHLESLFT	AITNFLLSLGI	HLNXNKTKR	MGA2TNF.
MGYGIGSWG				

Figure 17

Figure 18	: Patient	E HBsAg			
		20 WNSSTFHQA	30 LLDPRVRGLY	40 FPAGGSSSGT	50 VNP
-	•	70 APNMESTTS		90 AGFFLLTRIL	100 TIP
11 QSLDSWWTS		-		140 PICPGYRWMC	150 LRR
16 FIIFLFILL				190 STGPCKTCTT	200 PAQ
21 GTSMFPSCC		20 TCIPIPSSW		240 SVRFSWLSLL	250 VPF
26 VQWFVGLSF			280 NILNPFLPLL	290 PIFFCLWVYI	300 *TX
TKPNVGA					

Figure 18

Figure 19: Patie	ent F: nt	sequence		
10	20	30	40	50
CCAATCGGCAGTCA				
60			90	100
GACAGTCATCCTCA	GGCCATGCAG	TGGAACTCCA	GCACATTCCA	CCAAGC
110		130		150
TCTGCTAGATCCCA	SAGTGAGGGG	CCTATACTTT	CCTGCTGGTG	GCTCCA
160	170	180	190	200
GTTCCGGAACAGTA				
GIICCGGAACAGIA	MCCCIGIIC	CONCINCIOC	01010001111	
210	220	230	240	250
ATCTTCTCGAGGAC'				CACATC
260	270	280	290	300
AGGATTCCTAGGAC	CCCTGCTCGT	GTTACAGGCG	GGGTTTTTCI	TGTTGA
310	320	000		350
CAAGAATCCTCACA	ATACCACAGA	GTCTAGACTC	GTGGTGGACT	TCTCTC
260	270	200	200	400
<del>-</del>		380		
AATTTTCTAGGGGG	AGCACCCACG	TGTCCTGGCC	AAAAIIIGCA	3610000
410	420	430	440	450
AACCTCCAATCACT				
	00			
460	470	480	490	500
GCTGGATGTGTCTG	CGGCGTTTTA	TCATCTTCCT	CTTCATCCTC	CTGCTA
510		530		550
TGCCTCATCTTCTT	GTTGGTTCTT	CTGGACTACC	AAGGTATGTI	GCCCGT
	530		500	<b>CO</b> 0
		580		
TTGTCCTCTACTTC	CAGGAACATC	AACTACCAGC	ACGGGACCAT	GCAAGA
610	620	630	640	650
CCTGCACGACTCCT				
501501.001.01				

Figure 19

660	670	680	690	700		
ACAAAACCTTCGGACG	GAAATTGCAC'	TTGTATTCCC	ATCCCATCAT	CTTG		
710		730		750		
GGCTTTCGCAAGATTC	CTATGGGAGT	GGGCCTCAGT	CCGTTTCTCC'	rggc		
760	770	780	790	800		
TCAGTTTACTAGTGCC	ATTTGTTCAG	TGGTTCGTAG	GGCTTTCCCC	CACT		
810	820	830	840	850		
GTTTGGCTTTCAGTTA	TATGGATGAT	GTGGTATTGG	GGGCCAAGTC'	rgta		
•						
860	870	880	890	900		
CAACATCTTGAATCCC	TTTTTACCGC'	TGTTACCAAT	TTTCTTTTGT	CTTT		
910	920	930	940	950		
GGGTATACATTTAAAC	CCTACTAAAA	CTAAACGTTG	GGGCTACTCC	CTTA		
960	970	980				
ACTTCATGGGATATGTAATTGGAAGTTGGGGTACCTTG						

Figure 19 continued

Figure 20 Patient F Pol Amino acid sequence

EDWGPCTE	10	20	30	40	50
	YGEHHIRIE	PRTPARVTGGV	FLVDKNPHNT	TESRLVVDFS	SQFS
RGSTHVSW	60	70	80	90	100
	PKFAVPNL(	QSLTNLLSSNL	SWLSLDVSAA	Yeyhlplhpaa	MPH
	.10	120	130	140	150
	.pryvarls:	STSRNINYQHG	TMQDLHDSCS	SRNLYVSLLLI	LYKT
	.60	170	180	190	200
	'SHPIILGF	RKIPMGVGLSE	FLLAQFTSAI	CSVVRRAFPI	ICLA
	210	220	230	240	250
	VLGAKSVQHI	LESLFTAVTNE	LLSLGIHLNE	PTKTKRWGYSI	LNFM
GYVIGSWO	<b>;</b>				

Figure 20



Figure 21 Patient F HBsAg seq

	10	20	30	40	50		
MESTTSGFI	LGPLLVLQA	GFFLLTRILT	I PQSLDSWWI	SLNFLGGAPT	CPG		
			•				
1	60	70	80	90	100		
QNLQSPTSNHSPTSCPPICPGYRWMCLRRFIIFLFILLLCLIFLLVLLDY							
	- •		130		150		
QGMLPVCPLLPGTSTTSTGPCKTCTTPAQGTSMFPSCCCTKPSDGNCTCI							
_			180		200		
PIPSSWAFARFLWEWASVRFSWLSLLVPFVQWFVGLSPTVWLSVIWMMWY							
2:	10	220					
WGPSLYNILNPFLPLLPIFFCLWVYI							

Figure 22: Patient G ; HBV nt

	•					
10	20	30	40	50		
TCCGCCTCCTGCCT	CCACCAATC	GCCAGTCAGGA	AGGCAACCTA	ACCCCGC		
.000000						
60	70	80	90	100		
TCTCTCCACCTTTG	AGAGACACT	CATCCTCAGGC	CGTGCAGTG	BAACTCC		
.0.0.00.00						
110	120	130	140	150		
ACAACCTTCCACCA				CTATCT		
11012100110011						
160	170	180	190	200		
CCCTGCTGGTGGCT	CCAGTTCAG	GAACAGCAAAC	CCTGTTCCG	ACTACTG		
00010010010010						
210	220	230	240	250		
CCTCTCGCTTATCG						
001010001111100						
260	270	280	290	300		
ATGGAGAACATCAC						
AI COMOMICA COM	01.0001					
310	320	330	340	350		
GGGGTTTTTCTTGT						
99991111101101	1 ONCENTOR II	1001011111	.00001.01.01			
360	370	380	390	400		
CGTGGTGGACTTCT	こっし ひんせんしん	CTACCCCCAA	тассетете:			
CGIGGIGGACIICI	0101101111	01110000001111				
410	420	430	440	450		
CAAAATTCGCGGTC						
CAAAATTCGCGGTC	CCCAACCIC	Criti Crici Cri	JOIN 1001 001			
460	470	480	490	500		
GACTTGTCCTGGTT.	ATCCCTCCA	TCTATCTCCCC	CGTTTTATC	ATATTCC		
GACTIGICCIGGII	AICGCIGGA	10111101000				
510	520	530	540	550		
TCTTCATCCTGCTG	320 CTATCCCTC	₯₼₵₼₼₵₼₼₵₼	rectrotrotr	GGACTAT		
ICTICATOCIGCIG	CINIGCCIC	ALOT TOTTOL	.001101101			
560	570	580	590	600		
CAAGGTATGTTGCC	ン / O CCTTTTCTCC	™CTATUTE & TOTAL				
CAAGGIAIGIIGCC	CGIIIGICC	1017111110011	00:11 00 1 0:11	001100110		
610	620	630	640	650		
CACGGGAACATGCCGAACCTGCACGACTCCTGCTCAAGGAACCTCTATGT						
660	670	680	690	700		
ATCCCTCCTGTTGC						
WICCCICCIGIIGC	TGIVCCVVV	COLLOGOROGI	ormer roome			

Figure 22

710	720	730	740	750			
CCCATCCCATCATCTTGGGCTTTCGGAAAATTCCTATGGGAGTGGGCCTC							
760			790	800			
AGCCCGTTTCTCCTGGCTCAGTTTACTAGTGCCATTTGTTCAGTGGTTCG							
810	820	830	840	850			
TAGGGCTTTCCCCCACTGTTTGGCTTTCAGTTATATGGATGATGTGGTAT							
860	870	880	890	900			
TGGGGGCCAAGTCTGTACAGCATCTTGAGTCCCTTTTTTACCGCTGTTACC							
910	920	930	940	950			
AATTTTCTTTTGTCTTTGGGTATACATTTAACCCCTAACAAAACAAAGAG							
960	970	980	990	1000			
ATGGGGTTACTCTCTAAATTTTATGGGCTATGTCATTGGAAGTTATGGGT							
1010	1020	1030	1040				
CCTTGCCACAAGAACACATTATACTAAAAATCAAAGATTGTTT							

Figure 22 continued

Figure 23 Patient G HBV POL

10	20	30	40	50
EDWGPCAEHGE	CHHIRTPRTPSRV	TGGVFLVDKNP	HNTAESRLV	VDFSQFS
60	70.	80	90	100
RGNYRVSWPKE	'AVPNLQSLTNLI	SSDLSWLSLDV	SAAFYHIPL	НРААМРН
110	120	130	140	150
LLVGSSGLSRY	VARLSSNSRILN	HQHGNMPNLHD	SCSRNLYVS	LLLLYOT
160	170	180	190	200
FGRKLHLYSHP	IILGFRKIPMGV	GLSPFLLAQFT	SAICSVVRR	AFPHCLA
		-		
210	220	230	240	250
FSYMDDVVLGA	KSVQHLESLFTA	VTNFLLSLGIH:	LTPNKTKRWO	SYSLNFM
			•	
GYVIGSYG.				

Figure 23



Figure 24: Patient G HbsAg

_	0	.20	30	40	50
	GPLLVLQA	GFFLLTRILT	IPQSLDSWWI	SLSFLGGTT	VCLG
_	0	70	80	90	100
	HSPTSCPP	TCPGYRWMYL	RRFIIFLFII	LLLCLIFLLVI	LLDY
11 QGMLPVCPL	-		130 AQGTSMYPSO	140 CCCTKPSDGNO	150 CTCI
16	•	170	180	190	200
PIPSSWAFG		ARFSWLSLLV	PFVQWFVGLS	SPTVWLSVIW	MMWY
21 WGPSLYSIL		220 IFFCLWVYI			

Figure 24



#### Figure 25 Patient H nt seq

		30		50
CGCCTCCTGCCTCC	ACCAATCGC	CAGTCAGGAAG	GCAGCCGACC	CCACTG
	70		90	100
TCTCCACCTTTGAG	AGACACTCAI	CCTCAGGCCG	TGCAGTGGA	ACTCCAC
110		130		
AACCTTCCACCAAA	CTCTGCAAG	ATCCCAGAGTG	AGAGGCCIGI	ATTICC
160		180		
CTGCTGGTGGCTCC.	AGTTCAGGA <i>i</i>	ACAGTAAACCC	TGTTCCGACC	CACTGCC
210		230		250
TCTCCCTTATCGTC	AATCTTCTC	SAGGATTGGGG	ACCCTGCGC'1	GAACAT
260		280		
GGAGAACATCACAT	CAGGATTCC	PAGGACCCCTT	CTCGTGTTAC	CAGGCGG
310		330		
GGTTTTTCTTGTTG.	ACAAGAATC	CTCACAATACC	GCAGAGTCT?	AGACTCG
360		380		400
TGGTGGACTTCTCT	CAGTTTTCT	AGGGGAAACCA	CCGTGTGTCT	TGGCCA
		430		450
AAATTCGCAGTCCC	CAACCTCCAI	ATCACTCACCA	ACCTCCTGT	CCTCCAA
460	470 .	480	490	500
CTTGTCCTGGTTAT	CGCTGGATG!	rgtctgcggc	TTTTATCAT!	ATTCCTC
510		530		550
TTCATCCTGCTGCT	ATGCCTCAT(	CTTCTTGTTGG	STTCTTCTGG?	ACTATCA
560		580		600
AGGTATGTTGCCCG	TTTGTCCTC	PAATTCCAGG <i>F</i>	ATCCTCAACC	ACCAGCA
		630		650
CGGGACCATGCCGA	ACCTGCACG	ACTCCTGCTCF	AGGAACCTC	PATGTAT.
	670			700
CCCTCCTGTTGCTG	TACCAAACC!	TTCGGACGGA <i>F</i>	ATTGCACCT(	STATTCC

Figure 25

7	10	720	730	740	750
CATCCCAT	CATCTTGG	GCTTTCGCAA!	AATTCCTATG(	GGAGTGGGG	CTCAG
	60				800
CCCGTTTC	TCATGGCT	CAGTTTACTAC	GTGCCATTTG!	rtcagtggt1	CGTA
. 8	10	820	830	840	850
GGGCTTTC	CCCCACTG	rttggctttc <i>i</i>	AGTTATGTGG!	ATGATGTGGT	TATTG
8	60	870	880	890	900
			GTCCCTTTTT	ACCGCTGTT	ACCAA
g	10	920	930	940	950
TTTTCTTTTGTCTTTGGGTATACATTTAAACCCTAACAAAACGAAAAGAT					
g	60	970	980	990	1000
GGGGTTAC	TCTTTAAA'		PATGTTATTG(		GTCC
10	10	1020			
TTGCCACA	AGAACACA'	TCGTACAAAA	A		

Figure 25 continued



Figure 26: Patient H HBV pol

10	20	30	40	50
EDWGPCAEHGEHHI	RIPRTPSRV	TGGVFLVDKNP	HNTAESRLVV	DFSQFS
		•		
60	70	80	90	100
RGNHRVSWPKFAVP	NLQSLTNLL	SSNLSWLSLDV	SAAFYHIPL	IPAAMPH
110	120	130	140	150
LLVGSSGLSRYVAR	LSSNSRILN	HQHGTMPNLHD	SCSRNLYVSI	LLLYQT
	•			
160	170	180	190	200
FGRKLHLYSHPIIL	GFRKIPMGV	GLSPFLMAQFT	SAICSVVRR	AFPHCLA
. 210	220	230	240	250
FSYVDDVVLGAKSV:	SHLESLFTA	VTNFLLSLGIH	LNPNKTKRWO	GYSLNFM
260				
GYVIGCYGSLPQEH				

Figure 26

Figure 27: Patient H HBsAg

10	20	30	40	50	
MENITSGFLG	PLLVLQAGFF	LLTRILTIPQ	SLDSWWTSLSI	FLGETTVCLG	
60	70	80	90	100	
QNSQSPTSNH	SPTSCPPTCP	GYRWMCLRRF	IIFLFILLLC	LIFLLVLLDY	
110	120	130	140	150	
QGMLPVCPLI:	PGSSTTSTGP	CRTCTTPAQG	TSMYPSCCCTI	KPSDGNCTCI	
160	170	180	190	200	
PIPSSWAFAKFLWEWGSARFSWLSLLVPFVQWFVGLSPTVWLSVMWMMWY					
210	220				
WGPSLYRILS					
MAESTIVITIS	ELMENDETER	CTMATT			

Figure 27

#### Figure 28 Patient I HBV nt seq

10	20	30	40	50
CAACTTGTCCTGG	PTATEGETGG	ATGTGTCTGCG	GCGIIIIAIC	AIAIIC
60	70	80	90	100
CTCTTCATCCTGC'	PGCTATGCCT(	CATCTTCTTGT	TGGTTCTTCI	GGACTA
110	100	120	140	150
110	120	130		
TCGAGGTATGTTG	CCCGTTTGTCC	CTCTACTTCCA	AGGATCTTCAP	ACCACCA
160	170	180	190	200
GCACGGGTCCATG			יייר א א ככא א כר	PTATO
GCACGGGICCAIG	SAGAACCIGCA	aconcicció.	JI CHI IOCIII IOC	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
			0.40	252
210		230	240	250
TATCCCTCATGTT	<b>GCTGTACCAA</b> ?	ACCTTCGGAC(	<b>GAAATTGCA</b> C	CTGTAT
260	270	280	290 ·	300
TCCCATCCCATCA				
TCCCATCCCATCA	TCCTGGGCTT.	I CGGAAAAI I (	CINIGGONG	GGGCCI
310	320	330	340	350
CAGCCCGTTTCTC	ATGGCTCAGT	TTACTAGTGC	CATTTGTTCAC	STGGTTC
000000				
260	270	380	390	400
360				
GTAGGGCTTTCCC	CCATTGTTTG	GCTTTCAGTT	ATGTGGATGAT	rGTGGTA
•				
410	420	430	440	450
TTGGGGGCCAAGT				CTGTTAC
11666666664461	CIGINICOCK.	10110110100	0111111000	
		400	400	500
460	470	480	490	500
CAATTTTCTTTTG	TCTCTGGGTA:	TACATTTAAA	CCCTCACAAA	ACAAAAA
510	520	530	540	550
GATGGGGTTACTC				татссс
GWIGGGGIIWCIC	TITOMITIC	11100011110		
560				
TCTTTGCCAC			•	

Figure 29 Patient I HBV pol

10 20 30 40 50

NLSWLSLDVSAAFYHIPLHPAAMPHLLVGSSGLSRYVARLSSTSRIFNHQ

60 70 80 90 100

HGSMQNLHDSCSRNLYVSLMLLYQTFGRKLHLYSHPIILGFRKIPMGVGL

110 120 130 140 150

SPFLMAQFTSAICSVVRRAFPHCLAFSYVDDVVLGAKSVSHLESLFTAVT

160 170 180

NFLLSLGIHLNPHKTKRWGYSLHFMGYVIGCYGSLP

Figure 29

Figure 30 Patient I: HBsAg

50	40	30	20	10
FPFECSSIL2	LDYRGMLPVCP	PICTIETTAL	RRFIIFLFILI	TCPGYRWMCLE
_				
100	90	80	70	60
GKFLWEWAS	<b>TCIPIPSSWAF</b>	CTKPSDGNC'	AOGTSMYPSCO	TGPCRTCTTPA
			-	
150	140	130	120	110
(LSPFLPLLP	MWYWGPSLYRI	PIVWLSVMWM	PFVQWFVGLSE	ARFSWLSLLVI
		180	170	160
				IFFCLWVYI*

Figure 30

Figure 31 Patie	ent J HBV n	t seq		
10	20	30	40	50
CGCCTCCTCCTGCC	CTCCACCATCG	GCAGTCAGG	AAGAAAGCCTA	CTCCCA
60	70	80	90	100
TCTCTCCACCTCT				AACTCC
110		130		
AGCACATTCCACCA	AAGCTCTGCTA	GATCCCARA	GTGAGRGGCCT	ATACTT
1.00	170	100	100	, 200
160 TCCTGCTGGTGGC	፲ / U ኮርር ልርጥጥርርርር	TOU TOU	T DO ADODTTOTOTO	CTACTG
TCCTGCTGGTGGC.	CCAGIICCGG	- ANOMOTIME	0001011000	.011.010
210	220	230	240	250
CCTCTCCCATATC	GTCAATCTTCT	CGAGGACTG	GGGACCCTGCA	CCGAAT
				_0_
	270			
ATGGAGAGCACAA	CATCAGGATTC	CTAGGACCC	CTGCTCGTGTT	ACAGGC
21.0	320	330	340	350
310 GGGGTTTTTCTTG'				
GGGGTTTTCTTG	IIGACAAGAAI	CCICACA		
360	370	380	390	400
CGTGGTGGACTTC'			CACCCACGTGT	CCTGGC
				450
	420			
CAAAATTTGCAGT	CCCCAACCTCC	AATCACTCA	CCAACCTCTTG	TCCTCC
460	470	480	490	500
AATTTGTCCTGGT				
111111101001001				
510			540	
TCTTCATCCTGCT	GCTATGCCTCA	TCTTCTTGT	KGGTTCTTCTG	GACTAC
		<b>500</b>	500	600
	570			
CAAGGTATGTTGC	CCGTTTGTCCT	CTACTTCCA	GGAACA I CAAC	IACCAG
610	620	630	640	650
CACGGGACCATGC				
•			_	
	670			
TTCCCTCTTGTTG	CTGTACAAAAC	CTTCGGACG	GAAATTGCACT	TGTATT

Figure 31

710	720	730	740	750
CCCATCCCATCATC'		CGCAAGATTCC	TATGGGAGT	GGCCTC
			790	800
AGTCCGTTTCTCCT	GGCTCAGTT	TACTAGTGCCA	TTTGTTCAGI	rGGTTCG
		•		
810	820		840	850
TAGGGCTTTCCCCC	ACTGTTTGG	CTTTCAGTTAT	'ATGGATGAT(	STGGTAT
	870		890	900
TGGGGGCCAAGTCT	GTACAACAI	CTTGAATCCCT	TTTTACCGC	TGTTACC
910	920	930	940	950
AATTTTCTTTTGTC	TTTGGGTAT	ACATTTAAACC	CTACTAAAA	CTAAACG
960	970	980	990	1000
TTGGGGCTACTCCC	TTAACTTC	TGGGATATGTA	ATTGGAAGT	rggggta
1010	1020			
CCTTACCACAGGAA	CATATTGT	ACACAAA		

Figure 31 continued

Figure 32 Patient J HBV pol

EDWGPCT		20 PRTPARVTGG	30 VELVDKNPHN	40 FTESRLVVDFS	50 SQFS
RGSTHVS	60 WPKFAVPNLÇ	70 QSLTNLLSSN	• •	90 AFYHLPLHPA	100 AMPH
		120 STSRNINYQHO		140 SRNLYVSLLLI	150 LYKT
			180 PFLLAQFTSA:	190 ICSVVRRAFPI	200 ICLA
		220 LESLFTAVTN		240 PTKTKRWGYSI	250 LNFM
	260 GTLPQEHIVI	łK			

Figure 32

#### Figure 33. Patient J HBsAg

10	20	30	40	50
MESTTSGFLGE	PLLVLQAGFFLL	TRILTIPQSLD	SWWTSLNFLGG	APTCPG
60	70	80	90	100
QNLQSPTSNHS	SPTSCPPICPGY	RWMCLRRFIIF	LFILLLCLIFL	XVLLDY
110	120	130	140	150
QGMLPVCPLLE	PGTSTTSTGPCK	TCTIPAQGTSM	FPSCCCTKPSD	GNCTCI
160	170	180	190	200
PIPSSWAFARE	TLWEWASVRFSW	LSLLVPFVQWF	VGLSPĮTVWLSV	IWMMWY
210 WGPSLYNILNE	220 PFLPLLPIFFCL	WVYI		

Figure 33

Figure 34 Patient K HBV nt

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	10	20	30	40	50
CTCCT		CACCAATCGGC	AGTCAGGAAG	BACAGCCTACA	CCCATC
	60	70	80	90	100
TCTCC	ACCTCTAA	GAGACAGTCAT	CCTCAGGCCA	<b>ATGCAGTGGAP</b>	CTCCAG
•					
	110	120	130	140	150
CACAT	TCCACCAA	GCTCTGCTAGA	TCCCAGAGT	SAGGGGCCTAT	ACTTTC
	160	170	180	190	200
CTGCT		CAGTTCAGGAA	- · ·		CACTGCC
	210	220	230	240	250

260	270	280	290	300		
GGAGAGCACCACATCAGGATTCCTAGGACCCCTGCTCGTGTTACAGGCGG						

TCTCCCATATCGTCAATCTTCTCGAGGACTGGGGACCCTGCACCGAATAT

310	320	330	340	350	
GGTTTTTCTTGTTGACAAGAATCCTCACAATACCACAGAGTCTAGACTCG					

	360	370	380	390	400
TGGTGG	ACTTCTC	TCAATTTTCTA	GGGGGAGCA	CCACGTGTC	CTGGCCA

410	420	430	440	450	
AAATTTGCAGTCCCCAACCTCCAATCACTCACCAACCTCTTGTCCTCCAA					

460	470	480	490	500		
TTTGTCCTGGTTATCGCTGGATGTGTCTGCGGCGTTTTATCATCTTCCTC						

510	520	530	540	550		
TTCATCCTGCTGCTATGCCTCATCTTCTTGTTGGTTCTTCTGGACTACCA						

560	570	580	· 590	600			
AGGTATGTTGCCCGTTTGTCCTCTACTTCCAGGAACATCAACTACCAGCA							

			4			
610	620	630- ; ·	640	650		
CGGGACCATGCAAGACCTGCACGATTCCTGCTCAAGGAACCTCTATGTTT						

660	670	680	690	700		
CCCTCTTGTTGCTGTACAAAACCTTCGGACGGAAATTGCACTTGTATTCC						

## Figure 34

## . 45 of 51

•	710	720	730	740	750
CATCC	CATCATCTT	GGGCTTTCG	CAAGATTCCTA	TGGGAGTGG	SCCTCAG
	760			790	800
TCCGTT	TTCTCCTGG	CTCAGTTTA	CTAGTGCCATT	TGTTCAGTG	GTTCGTA
	810 <sup>-</sup>	820	830	840	850
GGGCT	TCCCCCAC	TGTTTGGCT'	TTCAGTTATAT	GGATGATGT	GGTATTG
	860	870	880		900
GGGGC	CAAGTCTGT	ACAACATCT	TGAATCCCTTT	TTACCGCTG	TTACCAA
			930		950
TTTTC	PTTTGTCTI	TGGGTATAC	ATTTAAACCCI	rctaaaacc	AAACGTT
eccemi			980 GGATATGTAA1		1000
GGGGT.	raciccum	AACTICATG	GGATATGIAAI	LIGANDIIG	
TTACC		1020 ATATTGTACA	1030 CAAAATÇAAAC	CA ·	

Figure 34 continued

Figure 35 Patient K HBV pol

	10	20		40	50
SSCLHQS	AVRKTAYTHI	LSTSKRQSSSC	SHAVELQHIP	PSSARSQSEGI	?ILS
	60	70	80	90	100
CWWLQFR				HIRIPRTPAR	
	110	120	130	140	150
VFLVDKN	PHNTTESRLY	/VDFSQFSRG	STHVSWPKFAV	/PNLQSLTNL	LSSN
	160	170			200
LSWLSLD	VSAAFYHLPI	LHPAAMPHLL	VGSSGLPRYVI	ARLSSTSRNII	HQY <i>I</i>
			230		250
GTMQDLH	DSCSRNLYV	SLLLLYKTFGI	RKLHLYSHPI:	ILGFRKIPMG	VGLS
			280		300
PFLLAQF	TSAICSVVRI	RAFPHCLAFS'	YMDDVVLGAK	SVQHLESLFT	AVTN
	212	222	220	240	
	310	320		340	
FLLSLGI	HLNPXKTKR	WGYSLNFMGY'	VIGSWGTLPQ	EHIVHKIK	

Figure 35

Figure 36 Patient K HbsAg

10 PPASTNRQS	) 20 GRQPTPISPPL			50 DPRVRGLYFP
60	70		90	100
AGGSSSGTVN	PVPTTASPIS		PNMESTTSGF	LGPLLVLQAG
110	120		140	150
FFLLTRILTI	PQSLDSWWTS		GQNLQSPTSI	NHSPTSCPPI
160	170	180	190	200
CPGYRWMCLR	RFIIFLFILL	LCLIFLLVLL	YQGMLPVCPI	LLPGTSTTST
210	220	230		250
GPCKTCTIPA	QGTSMFPSCC	CTKPSDGNCTO		ARFLWEWASV
260 RFSWLSLLVP	270 FVQWFVGLSP1	280 TVWLSVIWMMW	290 YYWGPSLYNII	300 NPFLPLLPI
310	320	330	340	'KSN
FFCLWVYI*T	LLKPNVGVTPI	LTSWDM*LEVG	VPYHRNILYT	

Figure 36

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Figure	37	Patient	L	HBV	nt
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. Iguic of Luc				
10 CAGTCCGGAAGG	20		40	50 ACACTCA
CAGTCCGGAAGG	CAGCCIACICC	LIMICICONC	,CICIAAGOA	101101011
60			90	
TCCTCAGGCCAT	'GCAGTGGAACT(	CCACCACTTTC	CATCAAACT	CTTCAAG
110	120			.150
ATCCCAGAGTCA	GGGCTCTGTAC!	TTTCCTGCTG6	TGGCTCCAG	rtcagga
160	170	180	190	200
ACAGTGAGCCCT	'GCTCAGAATAC'	TGCCTCTGCCA	TATCGTCAA	CTTCTC
210	220	230	240	250
GAAGACTGGGGA	ACCCTGTACCGA	ACATGGAGAA	CATCGCATCA	GGACTCC
0.11.01.01.000.	.0001011100011			
260	270	280	290	300
	CTCGCGTTACAG	<b>്ട</b> ്ട	rCTCGTTGAC	
TAGGACCCCIGC	,1CGCG11ACAG	GCGGGG1111.	.01001100.	
210	320	330	340	350
CTCACAATACCA	ACAGAGTCTAGA	CICGIGGIGG	ACTICICION	AIIIICI
		000	200	400
		380		
AGGGGGAACAC	CCGTGTGTCTTG	GCCAAAATTC	<b>SCAGTCCCAA</b>	ATCTCCA
		430		450
GTCACTCACCA	ACTTGTTGTCCT	CCAATTTGTC	CTGGTTATCG	CTGGATG
		480		
TGTCTGCGGCG:	TTTTATCATCTT	CCTCTGCATC	CTGCTGCTAT	GCCTCAT
510	520	530	540	550
СттСттсттСС	TTCTTCTGGACT	ATCAAGGTAT	GTTGCCCGTT	TGTCCTC
0110110110				
560	570	580	590	600
	TCATCAACCACC			
INDITIONS OF				
610	620	630	640	650
	AGGAACCTCTAT			
ACTOUTGUTUA	NGGMACCICIAI	GITICCCICA	TOLIGOIGIA	

Figure 37

660	670	680	690	700
TACGGACGGAAACT	GCACCTG1	FATTCCCATCCCA	TCATCTTGG	GCTTTCG
710	720	730	740	750
CAAAATACCTATGG	GAGTGGGC	CCTCAGTCCGTTT	CTCTTGGCT	CAGTTTA
760	770	780	790	800
CTAGTGCCGTTTGTT	CAGTGGT	TCGTAGGGCTTT	CCCCCACTG	CTGGCT
810	820	830	840	850
TTCAGTTATATGGAT	GATGTGG	TATTGGGGGCCA	AGTCTGTACA	AACATCT
860	870	880	890	900
TGAGTCCCTTTATGC	CGCTGTT	ACCAATTTTCTT	TTGTCTTTGG	GTATAC
910	920	930	940	950
ATTTAAACCCTCACA	AAACAAA	AAGATGGGGATAT	TCCCTTCAA	TTCATG
960 GGATATGTAATTGGG	970 GGTTGGG	980 GCTCCTTG		

Figure 37 continued

Figure 38. Patient L Pol

10 EDWGPCTEHGE	20 CHRIRTPRTPA	30 RVTGGVFLVI	40 OKNPHNTTESP	50 LVVDFSQFS
60	70	. 80	90	100
RGNTRVSWPKE	PAVPNLQSLTN	LLSSNLSWLS	SLDVSAAFYHĮ	PLHPAAMPH
110	120	130	140	150
LLVGSSGLSRY	VARLSSNSRI	INHQHRTMQN	NLHDSCSRNLY	VSLMLLYKT
160	170	180	190	200
YGRKLHLYSHE	PIILGFRKIPM	GVGLSPFLLA	OFTSAVCSVV	RRAFPHCLA
			_	
210	220	230	240	250
FSYMDDVVLGA	KSVQHLESLY.	AAVTNFLLSI	GIHLNPHKTK	RWGYSLQFM
	-			
GYVIGGWG				

Figure 38

## Figure 39 Patient L HBsAg

10	0	20	30	40	50
MENIASGLL	GPLLALQAG	FFSLTKILT	IPQSLDSWWT	SLNFLGGTPV	CLG
60	0	70 .	80	90	100
QNSQSQISSI	HSPTCCPPI	CPGYRWMCL	RRFIIFLCIL	LLCLIFLLVL	LDY
110	0 1	.20	130	140	150
QGMLPVCPL:	IPGSSTTST	GPCRTCTTP.	AQGTSMFPSC	CCTKPTDGNC	TCI
	•				
160	0 1	.70	180	190	200
PIPSSWAFA	KYLWEWASV	RFSWLSLLV	PFVQWFVGLS	PTVWLSVIWM	YWM
210	0 2	20			
WGPSLYNIL					

Figure 39

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